

WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification 4: A61K 39/005, 39/04, 39/40 C12N 15/00, 1/00, C12P 21/00 G01N 33/53, A61K 39/395

(11) International Publication Number:

WO 90/02564

A1

(43) International Publication Date:

22 March 1990 (22.03.90)

(21) International Application Number: .

PCT/US89/03955

(22) International Filing Date:

12 September 1989 (12.09.89)

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(30) Priority data:

243,474

12 September 1988 (12.09.88) US

(81) Designated States: AT (European patent), BE (European patent), CH (European patent), DE (European patent), FR (European patent), GB (European patent), IT (European patent), LU (European patent), NL (European patent), SE (European patent), US.

(60) Parent Application or Grant (63) Related by Continuation

US

243,474 (CIP)

Filed on

12 September 1988 (12.09.88)

Published

With international search report.

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CA 94080 (US).

Before the expiration of the time limit for amending the claims and to be republished in the event of the receipt of amendments.

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(54) Title: VACCINE DIAGNOSTIC EMPLOYING PROTEINS HOMOLOGOUS TO HEAT SHOCK PROTEINS OF TRYPANOSOMA CRUZI

(57) Abstract

This invention relates to vaccines and diagnostics and more particularly to vaccines and diagnostics which employ proteins and/or fragments and/or derivatives thereof having homology to heat shock proteins of Trypanosoma cruzi.

BNSDOCID: <WO 9002564A1 1 >

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VACCINE DIAGNOSTIC EMPLOYING PROTEINS HOMOLOGOUS TO HEAT SHOCK PROTEINS OF TRYPANOSOMA CRUZI

BACKGROUND OF THE INVENTION

This invention relates to vaccines and diagnostics and more particularly to vaccines and diagnostics which employ proteins and/or fragments and/or derivatives thereof having homology to heat shock proteins of Trypanosoma cruzi.

Heat shock proteins, sometimes referred to as stress proteins, have been found in a wide variety of cells, and have been generally described in an article written by Tissieres on pages 419 through 429 of "Heat Shock from Bacteria to Man" (Cold Spring Harbour Laboratory, 1982).

DESCRIPTION OF THE FIGURES

Figure 1 provides the gene and derived amino acid sequence for the Hsp70 antigen of T. cruzi.

Figure 2 provides an alignment of heat shock
proteins from a variety of organisms: 1. M. hyopneumoniae, 2. Bacillus megaterium, 3. Escherichia
coli, 4. T. cruzi, 5. T. cruzi, 6. Rat, 7. Xenopus
laevis 8. human, 9. chicken, 10. Zea mays, 11. Serratia
marcescens.

25 Figure 3 provides a restriction map of pMYCO16 containing the full length gene for the Hsp70 antigen of M. hyopneumoniae.

Figure 4 provides an intermediate plasmid for the expression of the Hsp70 antigen of M.

30 hyopneumoniae.

Figure 5 provides the gene and derived amino acid sequence for the Hsp70 antigen of M. hyopneumoniae.

Figure 6 provides restriction map of pMYCO29
35 which is a low level expression plasmid containing the

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full length gene for the Hsp70 antigen of M. hyopneumoniae.

Figure 7 provides a restriction map of pMYCO31 which is a high level expression plasmid containing the full length gene for the Hsp70 antigen of M. hyopneumoniae.

Figure 8 provides a restriction map of pCAM101 containing the trpT176 gene.

Figure 9 provides a restriction map of pMYCO32 which is an expression plasmid containing the full length gene for the Hsp70 antigen of M. hyopneumoniae and the trpT176 gene.

Figure 10 provides a restriction map of pMGA4 which is an expression plasmid containing the full length gene for the Hsp70 antigen of M. gallisepticum.

Figure 11 provides the gene and derived amino acid sequence for the Hsp70 antigen of M. hyopneumoniae.

Figure 12 provides a restriction map of pMGA10 which is an expression plasmid containing the full length gene for the Hsp70 antigen of M. hyopneumoniae and the trpT176 gene.

SUMMARY OF THE INVENTION

against organisms which comprise a physiologically acceptable carrier with a protein which is capable of eliciting an antibody which recognizes at least one epitope of a native protein present in the organism, the native protein having at least 50% homology with a heat shock protein of T. cruzi. Processes for protecting a host against an organism are also disclosed which comprise administering an effective amount of a protein capable of eliciting an antibody which recognizes at least one epitope of a native protein protein present in the organism, the native protein

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having at least 50% homology with a T. cruzi heat shock protein.

Further disclosed are processes for determining an organism in a host which comprise contacting a sample derived from a host containing an organism or suspected of containing an organism with an antibody or antibody fragment which recognizes at least one epitope of a native protein present in the organism, the native protein having at least 50% homology with a heat shock protein of T. cruzi; and determining protein present in the organism bound to the antibody.

For such vaccines and processes, the native

protein referred to above may be derived from a species of Mycoplasma, Mycobacteria or Trypanosoma, provided that the native protein is not derived from Trypanosoma cruzi. Preferably, the native protein of Mycoplasma derivation is one selected from the group consisting of M. mycoides, M. bovis, M. bovigenitalium, M. bovoculi, M. bovirhinis, M. dispar, M. hyorhinis, M. hyosynoviae, M. hyopneumoniae, M. gallisepticum, M. pneumoniae, and M. synoviae, most preferably from M. hyopneumoniae and M. gallisepticum. The native protein of Mycobacteria derivation is preferably one selected from the group consisting of M. bovis, M. leprae, and M. tuberculosis.

The recombinant sequence of nucleic acid

DETAILED DESCRIPTION

Applicant has found that certain heat shock proteins and/or fragments and/or derivatives thereof may be employed in a vaccine to protect against an organism containing such heat shock protein.

encoding the heat shock proteins of M. hyopneumoniae

and M. gallisepticum is also disclosed.

Applicant has further found that certain heat 35 shock proteins and/or fragments or derivatives thereof, as well as antibodies produced in response

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to such heat shock proteins and/or fragments or derivatives thereof may be employed as a diagnostic for determining an organism containing such heat shock proteins.

Applicant has also found that certain DNA (RNA) sequences encoding for a heat shock protein of an organism may be employed as a diagnostic for determining the organism.

In accordance with the one aspect of the present invention, there is provided a vaccine for protecting against an organism which includes a heat shock protein wherein the vaccine includes a protein capable of eliciting an antibody which recognizes at least one epitope of a heat shock protein of the organism which heat shock protein of the organism which heat shock protein of the organism has at least 50% homology with a heat shock protein of Trypanosoma cruzi (T. cruzi).

In accordance with another aspect of the present invention, there is provided a process for protecting against a disease caused by an organism which includes a heat shock protein by administering to a host at least one protein capable of eliciting an antibody which recognizes at least one epitope of a heat shock protein of the organism which heat shock protein of the organism has at least 50% homology with a heat shock protein of Trypanosoma cruzi (T. cruzi).

The term that an antigen or protein has at least 50% homology with a heat shock protein of T. cruzi, as used herein, means that on a position by position basis, at least 50% of the amino acids of the heat shock protein of T. cruzi are also present in the antigen or protein.

More particularly, in a preferred embodiment
the heat shock protein or polypeptide of T. cruzi with
which an antigen or protein is to have at least 50%
homology is at least one of the T. cruzi heat shock

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proteins having a molecular weight of about 70 kD, or about 85 kD or about 65 kD, preferably the heat shock protein having a molecular weight of about 70 kD.

The T. cruzi heat shock protein having a molecular weight of about 70 kD may be prepared as described in Example 1. The amino acid and DNA sequence for the 70 kD protein is shown in Figure 1 of the drawings, with the 70 kD protein starting at base pair 25 and terminating at base pair 677.

The T. cruzi heat shock protein having a molecular weight of about 85 kD is described by Dragon et al. Molecular and Cellular Biology, Volume 7 No. 3 Pages 1271-75 (March 1987).

The protein which is present in the organism and which is at least 50% homologous to a T. cruzi heat shock protein will sometimes be referred to herein as the "homologous protein" or the "homologous heat shock protein".

The protein employed in formulating the vaccine for protection against an organism may be identical to a homologous protein present in the organism to be protected against, or may be a fragment or derivative of such homologous protein, provided that the protein which is used in the vaccine is capable of eliciting an antibody which recognizes at least one epitope of the homologous protein. For example, the protein employed in the vaccine may be only a portion of the homologous protein present in the organism or may have one or more amino acids which differ from the amino acids of the homologous protein in the organism or may be the homologous protein (or fragment or derivative thereof) fused to another protein.

The term "protein which is capable of eliciting an antibody which recognizes at least one epitope of a native protein present in the organism, said native protein having at least 50% homology with a heat shock protein of T. cruzi" (such protein present

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in the organism is what is sometimes referred to as the homologous protein) encompasses the homologous protein present in the organism or a fragment of such homologous protein or a derivative of such homologous protein or a fusion product of such homologous protein (or fragment or derivative thereof) with another protein. As should be apparent, the protein or proteins included in the vaccine may include more or less amino acids or amino acids different from the amino acids of the homologous protein present in the organism.

The protein or proteins employed in the vaccine may be identified and produced by recombinant techniques. More particularly, the DNA (or RNA) encoding for a T. cruzi heat shock protein is employed as a probe to identify DNA present in the organism against which protection is sought which has at least 50% homology with the DNA (RNA) encoding for a T. cruzi heat shock protein. The DNA of the organism having the requisite homology is sometimes referred to herein as the "homologous DNA".

The homologous DNA of the organism identified by such probe is employed to produce homologous protein of the organism by recombinant techniques. Thus, for example, the DNA encoding for the protein of Figure 1 may be suitably labeled, for example with ³²P, by procedures known in the art to thereby provide a probe for identifying DNA in the organism having at least 50% homology with the DNA sequence encoding for the protein of Figure 1.

Figure 2 presents an alignment of the amino acid sequences of Hsp70 proteins from a number of species. The amino acids are depicted by their single letter abbreviations. Stretches of sequence identical in all examined species were identified (denoted by upper case text in the consensus sequence depicted below the individual sequences). Several regions

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containing sequences at least six amino acids in length which were identical in all Hsp70 sequences. For example, between amino acid 138 and 209 of T. cruzi lie the sequences TVPAYF, RIINEPTA, and DLGGGTFD which are conserved in Hsp70 sequences. The DNA sequences which could encode these conserved sequences were determined. The 17-mer nucleotide sequences having low coding degeneracy serve as universal oligonucleotide probes for Hsp70 genes.

The probing conditions selected are such that hybrids are identified in which there is at least 50% homology between the selected DNA probe which encodes for a T. cruzi heat shock protein and the DNA being probed for in the organism. Such probing is done at relatively low stringency. Low stringency is achieved by known methods such as reduced temperature and increased salt concentrations (e.g., hybridizing at 37°C and 5-6 X standard salt-citrate buffer or 5-6X standard salt-EDTA-Tris buffer).

The selected homologous DNA of the organism may be included in any of a wide variety of vectors or plasmids for producing a protein to be employed in formulating a vaccine against the organism. Such vectors include chromosomal, nonchromosomal and synthetic DNA sequences; e.g., derivatives of SV40; bacterial plasmids; phage DNA's; yeast plasmids; vectors derived from combinations of plasmids and phage DNAs, viral DNA such as vaccinia, adenovirus, fowl pox, virus, pseudorabies, etc.

The appropriate DNA sequences may be inserted into the vector by a variety of procedures. In general, the DNA sequences are inserted into an appropriate restriction endonuclease site by procedures known in the art. Such procedures and others are deemed to be within the scope of those skilled in the art.

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The DNA sequences in the vector are operatively linked to an appropriate expression control sequence(s) (promoter) to direct mRNA synthesis. As representative examples of such promoters, there may be mentioned: LTR or SV40 promoter, the E. coli lac or trp, the phage lambda PL promoter and other promoters known to control expression of genes in prokaryotic and eukaryotic cells or their viruses.

The expression vector also contains a ribosome binding site for translation initiation and a transcription terminator. The vector may also include appropriate sequences for amplifying expression.

In addition, the expression vectors

15 preferably contain a gene to provide a phenotypic trait
for selection of transformed host cells such as
dihydrofolate reductase or neomycin resistance for
eukaryotic cell culture, or such as tetracycline or
ampicillin resistance in E. coli.

20 The vector containing the appropriate DNA sequences as hereinabove described, as well as an appropriate promoter or control sequence, may be employed to transform an appropriate host to permit the host to express the protein. As representative 25 examples of appropriate hosts, there may be mentioned: bacterial cells, such as E. coli, Salmonella typhimurium, fungal cells, such as yeast; animal cells such as CHO or Bowes melanoma; plant cells, etc. The selection of an appropriate host is 30 deemed to be within the scope of those skilled in the art from the teachings herein.

The expression vehicle including the appropriate DNA sequences for the protein to be expressed and the t-RNA inserted at the selected site may include a DNA or gene sequence which is not part of the gene coding for the protein. For example, the desired DNA sequence may be fused in the same reading

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frame to a DNA sequence which aids in expression or improves purification or permits increases in the immunonogenicity.

In employing recombinant techniques for producing the active protein, purifications, digestions, ligations and transformations may be accomplished as described in "Molecular Cloning: A Laboratory Manual" by Maniatis et al., Cole Spring Laboratory, 1982 ("Maniatis"). In addition, transformations may be accomplished by the procedure

transformations may be accomplished by the procedure of Cohen, PNAS, 69:2110 (1973).

When seeking to develop a vaccine, neutralizing or protective antibodies could be targeted toward discontinuous, conformation-dependent epitopes of the native antigen. One must therefore consider 15 whether the protein obtained from the recombinant expression system might have a three dimensional structure (conformation) which differs substantially from that of the original protein molecule in its 20 natural environment. Thus, depending on the immunogenic properties of the isolated proteins, one might need to renature it to restore the appropriate molecular conformation. Numerous methods for renaturation of proteins can be found in the scientific literature and include; 1) denaturation (unfolding) of 25 improperly folded proteins using agents such as alkali, chaotropic agent, organic solvents, and ionic detergents followed by a renaturation step achieved by dilution, dialysis, or pH adjustment to remove the denaturant, and 2) reconstitution of proteins into a 30 lipid bilayer or liposome to re-create a membrane like environment for the immunogenic protein.

The vaccine which includes a protein of the type hereinabove described may be employed in a vaccine for protecting against diseases caused by a wide variety of organisms. Table 1 provides representative examples of such organisms. Of particular interest are

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species of Trypanosoma, Mycoplasma and Mycobacteria. Trypanosoma and Mycoplasma heat shock proteins are described herein. Heat shock proteins for Mycobacteria are known. Young et al., P.N.A.S. (USA), 85:4267-4270 (1988).

A host may be protected against a disease caused by a certain organism by incorporating into the vaccine a protein which is capable of eliciting antibodies which are recognized by at least one epitope of a homologous protein of the organism. As hereinabove indicated the protein which is capable of eliciting such antibodies (hereinafter sometimes referred to as the active protein) may correspond to the homologous protein of the organism or may be a fragment or derivative thereof. As should be apparent, if the disease against which a host is to be protected is Chagas, which is caused by T.cruzi, the protein which is included in the vaccine would be one or more heat shock proteins of T. cruzi or a fragment or derivative thereof capable of eliciting antibodies which recognize an epitope of T. cruzi heat shock The host which is protected is dependent upon the organism against which protection is sought. general, the host is an animal (either a human or nonhuman animal) which is subject to a disease caused by the organism. Thus, for example if the organism against which protection is sought is one which is known to cause disease in man, then the vaccine including the active protein or proteins would be administered to a human host. If the organism is known to cause a disease in a nonhuman animal, then the vaccine including the active protein would be administered to a nonhuman animal.

In formulating a vaccine, the active protein is employed in the vaccine in an amount effective to provide protection against the disease caused by the organism against which protection is sought. In

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general, each dose of the vaccine contains at least 5 micrograms and preferably at least 100 micrograms of the active protein. In most cases, the vaccine does not include the active protein in an amount greater than 20 milligrams.

The term "protection" or "protecting" when used with respect to a vaccine means that the vaccine prevents the disease or reduces the severity of the disease.

The active protein is employed in conjunction with a physiologically acceptable vehicle to provide protection against the organism. As representative examples of suitable vaccines in carriers, there may be mentioned: mineral oil, alum, synthetic polymers, etc. Vehicles for vaccines are well known in the art and the selection of a suitable vehicle is deemed to be within the scope of those skilled in the art from the teachings herein. The selection of a suitable vehicle is also dependent upon the manner in which the vaccine is to be administered. The vaccine may be in the form of an injectable dose and may be administered intra-muscularly, intravenously, or by sub-cutaneous administration. It is also possible to administer the vaccine orally by mixing the active components with feed or water; providing a tablet form, etc.

Other means for administering the vaccine should be apparent to those skilled in the art from the teachings herein; accordingly, the scope of the invention is not limited to a particular delivery form.

It is to be understood that a vaccine may also be formulated by use of an antibody elicited in response to a homologous protein of the organism.

The protein and/or antibody used in the vaccine is essentially free of the organism; i.e., cellular matter.

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In accordance with another aspect of the present invention, there is provided a diagnostic kit and/or assay for determining an organism which employs in the assay and/or kit an antigen which is recognized by an antibody elicited by a protein of the organism which has at least 50% homology with a T. cruzi heat shock protein, as hereinabove described, i.e., a "homologous protein" of the organism.

The antigen employed as a diagnostic may be obtained or produced as hereinabove described with reference to the active protein included in the vaccine.

In accordance with yet a further aspect of the present invention, there is provided a diagnostic assay and/or reagent for determining an organism which includes and/or employs an antibody (or fragment thereof) which recognizes an antigen of the organism to be determined, which antigen of the organism has at least 50% homology with a heat shock protein of T. cruzi, as hereinabove described.

The antibody employed in the assay and/or assay kit may be either a polyclonal or monoclonal antibody elicited in response to a homologous protein. In particular, the antibody employed in the diagnostic assay and/or kit is elicited in response to a protein and/or fragment and/or derivative thereof having at least 50% homology with a heat shock protein of T. cruzi.

A diagnostic kit and/or assay for determining 30 an organism which includes a homologous protein may be formulated to determine such organism by a variety of procedure.

For example, the organism may be determined by a so-called sandwich assay kit or assay for determining the organism by determining in a sample (derived from a host containing or suspected of

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containing the organism) antibody elicited in response to a homologous protein of the organism. In this procedure, antigen of the type hereinabove described is contacted with the sample under conditions at which any of such antibody present in the sample is immunobound to the antigen, which antigen is preferably supported on a solid support.

Antibody bound to such antigen may then be determined by use of an appropriate tracer comprised of a ligand bound or recognized by such antibody labeled with a detectable marker or label. The ligand of the tracer may be, for example, an antibody bound by or recognized by the bound antibody.

The marker may be any one of a wide variety of labels (for example a radioactive label, an enzyme label, a chromogen label, etc.).

The techniques for forming such an assay and for providing a tracer are known in the art and no further details in this respect are deemed necessary for understanding the present invention.

For example, there may be employed a so-called ELISA sandwich assay format in which a plastic microtiter plate is coated with an antigen of the type described (one which is recognized by antibody elicited in response to homologous protein of the organism) and sample derived from a host suspected of containing the organism is incubated with the coated antigen. After appropriate washing, labeled immunoglobulin (antiglobulin to the host species which is suspected of containing the organism) labeled with a detectable enzyme (for example horseradish peroxidase or alkaline phosphatase) is incubated with the antibody bound by the coated antigen. After washing, an appropriate developer is added.

Alternatively, an agglutination assay may be employed in which case particles, such as polystyrene

antibody.

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beads, coated with the appropriate antigen is mixed with appropriate sample, and presence of antibody is detected by agglutination.

These and other procedures should be apparent to those skilled in the art.

In an alternative sandwich immunoassay format, an antibody of the type hereinabove described may be employed to directly determine a homologous heat shock antigen or protein of the organism to be determined. For example, a sample (derived from a host 10 .containing or suspected of containing the organism) is subjected to a sandwich assay by contacting the sample with an antibody (or fragment thereof) which recognizes the homologous heat shock antigen of the organism, which antibody is preferably supported on a solid 15 support. Such contacting is effected under conditions which will immunobind the homologous heat shock antigen (if present) to the antibody. Thereafter, bound antigen may be determined by use of a tracer comprised 20 of a ligand (which is bound by or recognizes the homologous antigen) labeled with a detectable marker Thus, for example, the tracer may be labeled antibody elicited in response to the homologous antigen of the organism. As hereinabove indicated, the 25 antibodies capable of recognizing a homologous protein of the organism may be a monoclonal and/or polyclonal

In this assay format, which employs an antibody which recognizes a homologous protein of the organism, markers (labels) and techniques, as hereinabove described and as known in the art, may also be employed.

The assay or reagent kit which employs antigen and/or antibody of the type hereinabove

described may be included in an appropriate reagent kit package. The package may include other materials

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useful in the assay, for example, tracer, buffers, standards, etc., in appropriate reagent containers.

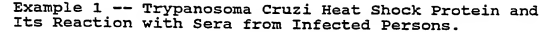
In accordance with another aspect of the present invention, there is provided an assay and/or reagent kit for determining the presence of an organism which includes or employs a DNA probe which encodes for a protein of the organism having at least 50% homology with a heat shock protein of T. cruzi as hereinabove described.

The DNA probe which is used may be all or a portion of the DNA which encodes for a homologous protein. If a portion of the DNA which encodes for a homologous protein is employed, such DNA portion should include a portion of the DNA which encodes for a variable region of the homologous protein.

Accordingly, the DNA probe is employed under conditions whereby hybridization is accomplished over at least a portion of the DNA which encodes for a variable region (preferably a hypervariable region) of the homologous protein.

The hydridization may be performed with a suitably labeled form of the DNA (for example ³²P, although other detectable labels, including non-radioactive labels may be used) in a procedure similar to the procedure for identifying DNA of the organism encoding for a protein having the requisite homology with a T. cruzi heat shock protein.

The present invention will be further described with respect to the following examples; however, the scope of the invention is not to be limited thereby. Unless otherwise indicated, all methods and abbreviations are well known in the art and are found in Maniatis. All references in this document are hereby incorporated by reference herein.



A. Growth and Isolation of Parasites

Trypanosoma cruzi, Peru strain, was used in 5 all experiments. Epimastigotes were grown at 28°C in modified HM (Warren, S. Parasitology, 46:529-539, 1960); 37 g/l brain heart infusion (Difco Lab., Detroit, MI), 2.5 mg/l hemin, 10% heat-inactivated fetal calf serum. Log phase cells were harvested by 10 centrifugation and washed twice with cold PSG (20 mM sodium phosphate, pH 7.4, 0.9% NaCl, 1.0% glucose). Culture form trypomastigotes were obtained from infected Va-13 cells as previously described. Sanderson et al., Parasitology, 80:153-162, (1980), and 15 Lanar and Manning, Mol. and Biochem., Parasitology, 11:119-131, (1984).

B. Isolation of DNA and RNA

Parasites were harvested from culture by centrifugation and washed several times with PSG (20 mM sodium phosphate, pH 7.4, 0.9% NaCl, 1.0% glucose). 20 Epimastigotes were resuspended at a concentration of 109/ml in PEG/EGTA buffer (20 mM Tris-HCl, pH 7.6, 25 mM EGTA, 50 mM MgCl, 25mM CaCl, 1.0% Triton-X100, and 4mM dithiothreitol), plus 250 u/ml of RNAS in (Promega Biotec, Madison, WI), incubated on ice for 20 min., 25 centrifuged at 8000 x g for 15 minutes at 4°C. The supernatant containing the RNA was phenol extracted 3 times, then extracted once with chloroformisoamyl alcohol (24:1) and ethanol precipitated. The pellet (nuclei and kinetoplasts) was resuspended at a 30 concentration of 109 parasite equivalents/ml in 10 mM Tris-HCl, pH 8.0, 50 mM EDTA, 0.1% SDS, 150 ug/ml Proteinase K (Boehringer- Mannheim, Indianapolis, IN) . and incubated at 65°C for 1 hour. After cooling to room temperature, the DNA was gently extracted with an 35 equal volume of phenol for 1 hour. This extraction

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was repeated once, and the aqueous phase was extracted with chloroform-isoamyl alcohol (24:1) once. The DNA was recovered by ethanol precipitation. The DNA pellet was gently redissolved in 10 mM Tris-HCl. pH 8.0. 1 mM EDTA and treated with 0.15 mg/ml DNAse-free RNAse A for 30 minutes at room temperature. After RNAse digestion the sample was extracted once with phenol, once with chloroformisoamyl alcohol, and then precipated with ethanol. The size of the DNA was determined to be greater than 20 kilobase pairs (kb) on agarose gels. Trypomastigote DNA and RNA was prepared in an identical manner except that the parasites were resuspended at a concentration of 5 x 109/ml.

C. Preparation of A+ mRNA

Poly A+ containing RNA was isolated by Oligo(dT)-cellulose chromatography (Aviv and Leder, J. Immunol., 127:855-859, 1972). Total RNA was loaded onto an oligo (dT)-cellulose column (Type 3, Collaborative Research, Lexington, MA) in 10 mM

Tris-HCl, pH 7.5, 1 mM EDTA, 0.2% SDS, 400 mM LiCl. RNA was eluted from the column at 40°C with 10 mM Tris-HCl, pH 7.5, 1 mM EDTA, 0.2% SDS.

D. Construction of the T. cruzi "Sau3a Partial" Genomic Library in Bacteriophage EMBL3

25 200 μg of T.cruzi epimastigote DNA was digested with the restriction endonuclease Sau3A (Boehringer-Mannheim, Indianapolis, IN) according to manufacturer's specifications. Aliquots of the reaction were removed at 1, 2.5, 5, 10, 20, 40 and 60 minutes. Upon removal each aliquot was diluted to 25 mM in EDTA and heated for 15 minutes at 68°C. The samples were pooled, the DNA was size fractionated over a Sephacryl S-1000 column (Pharmacia, Piscataway, NJ) in 200 mM Tris-HC1, pH 7.5, 100 mM NaCl, 1 mM EDTA.

35 Those fractions containing DNA in size from 5 kb to 20

kd were pooled, ethanol precipiated, and used for cloning. The lambda bacteriophage cloning vector EMBL3 (Frishauf et al., J. Mol. Biol., 170:827-842, 1983) was used. EMBL3 arms and GIGAPAK packaging system were purchased from Vector Cloning Systems (San Diego, CA) and used according to the manufacturer's instructions.

E. Hybridization-Selection/Translation

Specific T. cruzi RNAs were purified from total T. cruzi RNA using the technique of hybridization-selection/translation as described by 10 Riccardi et al., PNAS, 76:4927-4931, 1972. 25-50 ug of purified plasmid DNA was digested with an appropriate restriction endonuclease (to linearize the plasmid), the DNA was cleaned by phenol extraction and chloroform extraction and denatured by boiling for 10 minutes. 15 Following boiling, the DNA was quick-frozen, thawed, then spotted onto a 9mm diameter nitrocellulose filter. The filter was washed several times with 6XSSC, then air dried and baked for 2 hours at 80°C in vacuo. 20 hybridization, 100 μg of T. cruzi total RNA was reacted with the DNA containing filter in a solution containing 65% formamide, 0.01 M PIPES, pH 6.4, 0.4 M NaCl at 65°C for 3 hours. Following the hybridization reaction, the filter was washed 10 times with 1XSSC, 0.1% SDS at 60°C, 3 times with 0.002 M EDTA at 60°C, 25 and once with water at room temperature. specifically hybridized mRNA is eluted from the filter by boiling the filter in a small volume of water for two minutes, quick-freezing the solution, then ethanol precipitating the RNA. The purified RNA is resuspended 30 in water, then translated in an in vitro translation system (such as rabbit reticulocyte).

F. Immunoprecipitation Reactions

A 1:10 to 1:50 dilution of individual serum was prepared using the 10 mM Tris-HCl, pH7.5, 1% Nonidet

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P-40 (NP 40), 1 mM N-alpha-p-tosyl-L-Lysine chloromethyl ketone (TLCK), 1 mM phenyl methyl sulfonyl fluoride (PMSF), and 2.8 Kallikrein Inactivator Units (KIU)/ml aprotinin. The diluted serum was mixed with an equal volume of cell-free translation reaction mixture, and incubated overnight at 4°C. 10 µl of 10% protein-A-Sepharose (Pharmacia, Piscataway, NJ) was added and gently mixed for 1 hour at 4°C. The immune complexes were washed and analyzed on SDS-polyacrylamide gels as described in Dragon et al., Mol. and Biochem., Parasitology, 16:213-229, 1985.

G. Synthesis of cDNA

cDNA was synthesized by methods known to those of ordinary skill in the art. Briefly, 2 μ g of epimastigote or trypomastigote A+ mRNA was transcribed by the action of AMV reverse transcriptase as described by Ullrich et al., Science, 196:1313-1319, (1977) and Gubler, Gene, 25:263-269, (1983). Transcription was initiated at the 3' polyadenylated end of the mRNA using oligo(dt) as a primer. The second strand was copied using DNA polymerase I and RNAse H (Boehringer-Mannehim. Indianapolis, IN) and appropriate buffers.

Specifically, 2 µg of oligo-dT (12-18 nucleotides, Pharmacia Molecular Biology Division, Piscataway, NJ) was annealed to 2 micrograms of purified mRNA in the presence of 50 mM NaCl. The annealing reaction was heated to 90°C and then slowly cooled. For the reverse transcriptase reaction, deoxynucleosidetriphosphates (dATP, dTTP, dGTP and dCTP) were added to make a final concentration of 0.5 mM, along with 40 units of enzyme (Molecular Genetic Resources, Tampa, FL). The reverse transcriptase reaction buffer contained 15 mM Tris-HCl, pH 8.3, 21 mM KCl, 8 mM MgCl₂, 0.1 mM EDTA. and 30 mM beta-mercaptoethanol. This mixture was incubated at 42°C

for 45 minutes. The RNA-DNA duplex was extracted once with phenol chloroform and then precipitated with ethanol. The pelleted material was then resuspended in 100 microliter reaction mixture containing the following: 20 mM Tris-HCl pH 7.5, 5 mM MgCl₂, 100 mM KCl and 250 uM each dATP, dCTP, dTTP, dGTP.

RNAase H (100 units/ml Pharmacia Molecular Biology Division, Piscataway, NJ) and DNA Polymerase I -- Klenow fragment (50 units/ml Boehringer Mannheim, Indianapolis, IN) were added and the IO reaction was incubated at 12°C for 60 minutes. combined activities of these enzymes result in the displacement of the mRNA from the RNA-DNA duplex as the first cDNA strand is used as a template for 15 synthesis of the second cDNA strand. The reaction was stopped by the addition of EDTA to a final concentration of 10 mM and the DNA duplex was then extracted with phenol: chloroform and ethanol precipitated. The sequence of the reactions of DNA Polymerase I and RNAase H was predicted to yield cDNA 20 molecules which were blunt ended at both their 3' and 5' ends. A 3' blunt end is necessary for the subsequent cloning of the cDNA.

H. Construction of the cDNA Library

25 Briefly, the double stranded cDNA
preparations were digested with the restriction
endonucleases SacI and PvuII (New England Biolabs,
Beverly, MA) and ligated, using T4 DNA ligase, into the
SacI and SmaI sites of the plasmid pUC18 (Yanish-Perron
30 et al., Gene, 33:103-119, 1985). This mixture was used
to transform E. coli K12 strain JM83, selecting for
ampicillin resistance conferred by the introduction of
the pUC18 into the host cell. From 2 ug of mRNA
approximately 150 ng of cDNA were prepared which
yielded about 7000 ampicillin resistant clones.

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More specifically, the cDNA was resuspended in 100 microliters of sterile water. Approximately 50 ng was digested with SacI (5000 units/ml) and pVUII (12000 units/ml) in the presence of 6 mM Tris-HCl (pH and 6 mM beta-mercaptoethanol for 60 7.4) 6 mM MgCl2' minutes at 37°C.

The sample was then re-extracted with phenol: chloroform and ethanol precipitated. For the cloning step a pUC18 vector was used. The vector had been digested with SacI and SmaI. SmaI provided the blunt end site necessary for ligation of the 3' end of the cDNA. The ligation reaction was performed using 40 ng of vector DNA and 50 ng of cDNA. Ligation was done overnight at 12°C in a ligase buffer of 50 mM Tris-HC1 (pH 7.8), 10 mM MgC12, 20 mM dithiothreitol, 1.0 mM rATP using one unit of T4 DNA ligase.

The recombinant DNA molecules were then introduced into E. coli K-12 strain JM83 by transformation. The transformed bacteria were spread on agar plates containing the antibiotic ampicillin at a concentration of 50 micrograms/ml. Since the plasmid pUC18 contains the ampicillin resistance gene, only those bacteria which acquired a recombinant plasmid survived. These bacteria each grew and divided to form a bacterial colony. Each cell in the colony is a descendant of the original parental cell and contains the same recombinant plasmid. Using hybridization selection/translation and immunoprecipitation techniques to screen the cDNA library a clone was identified which contained nucleotide sequences corresponding to a 70 kd T. cruzi peptide.

I. Isolation of the full length 70 kd gene

The cDNA clone was used as a probe to screen the T. cruzi Sau3a partial genomic library as described by Maniatis et al. A lambda phage designated FG21 was identified which contained multiple copies of the 70 kD

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gene. A 2.4 kb Smal fragment was sub-cloned into pUC9 from FG 21. This subclone called pEG22 contained one full length copy of the 70 kD gene. The DNA sequence of PEG22 was determined. FG21, was sequenced and used to construct an expression plasmid to allow production of the 70 kd antigen in E. coli.

J. Expression of Cloned Genes in E. coli

Several systems are available in the laboratory for expressions of foreign genes in E. coli and other mammalian and bacterial tissue culture cell It is important to provide the cloned genes with an E. coli ribosome binding site for initiation of translation and a strong promotor to obtain sufficiently high levels of protein. obtaining "direct" expression of the protein is possible, it appears to be more efficient to produce the protein as a fusion protein, the amino.terminus of which is a small part of an E. coli protein containing signals for the initiation of protein synthesis. The amino terminus of B-lactamase and the amino terminus of B-galactosidase can make such fusion proteins [Hegpeth et al., Mol. Genet., 163:197-203 (1980) and Lingappa et al., PNAS, 81:456-460 (1984)]. These and other systems may be used to obtain expression of the cloned gene.

Sequencing analysis showed that the coding region of the 70 kd gene was flanked by an AhaIII site 30 base pairs upstream from the putative ATG start codon. An additional AhaIII site is located 367 base pairs following the TGA stop codon in the nucleotide sequence of FG21. Subsequently FG21 was digested with the restriction enzyme AhaIII. resulting DNA fragment was 2,341 base pairs long. was gel purified and cloned in the SmaI site of the 35 expression vector pUC9. The resulting plasmid, pFP70-47, was used to transform E. coli K12 SG936 bacteria.

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A sample of this recombinant bacteria has been placed on deposit with the American Type Culture Collection (12301 Parklawn Drive, Rockville, Maryland, USA) as ATCC number 67254. The culture was deposited on November 4, 1986. This strain, SG936/FP70-47 produces a 70 kd polypeptide which can react with chagasic sera. Expression of the entire protein, however, provides as many determinants as possible on the target antigen.

10 K. Antigen Production

The transformed E. coli are grown in liquid culture containing 50 micrograms per ml of ampicillin to enhance plasmid ability. Cultures are harvested at an OD of 2.0 measured at 550 nm. The cells are then pelleted and washed and lysed by freeze/thaw and sonication. A detergent extraction solubilizes most of the remaining polypeptides. The 70 kd expressed product, however, remains insoluble and is harvested by centrifugation. This insoluble "cement" is denatured in urea and subsequently diluted at a high pH and the pH is then adjusted back to neutral. During the renaturation process the antigen refolds and achieves that immunologically active conformation. The details of this procedure used are identical to those used to restore enzyme activity to recombinant chymosin as described by McCaman et al., J. Biotech., 12:117-191, (1985).

Example 2 -- 74.5 kda M. Hyo Antigen and Use As a Vaccine

30 A. Preparation of M. hyopneumoniae DNA

Strain P-57223 (obtained from Dr. Charles Armstrong, Purdue University) was grown in 1 liter of Friis medium to a density of approximately 10^9 to 10^{10} color changing units per ml. The cells were harvested by centrifugation and resuspended in 2 ml

phosphate buffered saline which brought the total volume to 3.25 ml. The suspension was then mixed with a solution consisting of 24.53 g cesium chloride dissolved in 19.75 ml 10 mM Tris pH 8.0 1 mM EDTA and 1.53 of 10 mg/ml ethidium bromide was added. 5 was mixed with a solution consisting of 3.87 g cesium chloride dissolved in 2.15 ml 10 mM Tris pH 8.0. 1 mM EDTA, 8.9% Sarkosyl. The resulting suspension was incubated at 65°C for 10 minutes to completely lyse the cells. The DNA was separated by equilibrium 10 buoyant density centrifugation in a Sorvall TV850 rotor at 43,000 rpm for 18 hours, and withdrawn with an 18 gauge needle. This DNA was subjected to two additional buoyant density centrifugations in a Sorvall TV865 rotor at 55,000 rpm for 7 and 18 hours 15 respectively, each time the band of genomic DNA being removed with an 18 gauge needle. The resulting DNA solution was extracted with cesium chloride saturated isopropanol, to remove ethidium bromide, and extensively dialyzed against 10 mM Tris pH 8.0, 1mM 20 EDTA, to remove the isopropanol and cesium chloride.

B. DNA Probing of M. hyopneumonia DNA

Plasmid pEG22, described in Example 1 is purified from E. coli by methods in the art, and labeled with ³²p by nick translation using DNA polymerase I.

pEG22 is used as a probe as follows:

Mycoplasma genomic DNA was digested with

EcoRI under the following conditions at 37°C for 2 hours.

- 114 microliters P-5722-3 DNA
 - 6 microliters H₂0
- 15 microliters 10X BRL-3 (Bethesda Research Labs)
- 15 microliters EcoRI (Bethesda Research Labs)

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67 microliters were mixed with 0.1% Bromphenol blue, glycerol, loaded onto a 1% agarose gel and electrophoresed until the blue color had migrated to within 1cm of gel end. The DNA was transferred to a nitrocellulose filter by Southern's technique. filter was hybridized to the DNA probe described above under conditions which allow hybridization in the absence of exact sequence identity.

Hybridization:

10 6 X NET

5 x Denhardts solution

2 X 106 counts per minute probe,

37°C for 18 hours

Wash:

15 6 X NET

0.1% SDS

3 times at room temperature,

1 time at 50°C

6 X NET

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20 1 M NaC1

90 mM Tris pH 7.6

6 mM EDTA

Southern blot analysis shows that the DNA probe hybridized to a specific EcoRI restriction endonuclease fragment of approximately 6 kB in length and thus include the antigen's gene.

C. Cloning the Gene by Hybridization

In order to identify the gene by hybridization to the pEG22 DNA probe, 200 micrograms of P-57223 DNA was digested with 120 units of EcoRI in a volume of 600 microliters. The digestion mixture was mixed with glycerol and xylene cyanol blue FF and electrophoresed on a 3.25% acrylamide gel.

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slices of approximately 0.5 cm were cut from the gel in the size range desired and electroeluted in 0.1% SDS, 0.5 X TBE buffer. The resulting DNA fractions were extracted with phenol/chloroform, ethanol precipitated, and each resuspended in 50 microliters of 10mM Tris pH 8.0, 1mM EDTA. By dot-blot analysis, (See Nuc. Acid Res. 7:1541-1552, 1979), fraction 4 was shown to contain the DNA fragment of interest.

To create a gene library enriched for the

desired fragment, 7 microliters of Fraction 4 was
ligated to EcoRI digested pUC9 with T4 ligase one-half
of the reaction was transformed into JM83 and plated on
X-gal plates where white colonies contain plasmids and
inserts. Plasmid DNA from 24 white colonies was
prepared and transferred to nitrocellulose by the
slot-blot modification of the dot-blot procedure and
probed with ³²P labeled pEG22.

Plasmid DNA preparations which hybridize to the DNA probe are subjected to EcoRI digest analysis to show that each plasmid contains the same size insert fragment, and most likely the same gene. A plasmid is selected for DNA sequence analysis which shows greater than 50% identity to pEG22.

D. Preparation of Genomic Library

A preparative digest of 200 μ g genomic DNA of Mycoplasma hyopneumoniae P-57223 was done using 200 units of EcoRI in a total volume of 1 ml and 250 μ l aliquots were removed at 6 min, 25 min, 42 min and 63 min.

30 The four preparative samples of partially digested Mycoplasma DNA were then combined (200 μ g) and loaded onto an exponential sucrose gradient. The gradient was centrifuged in a Sorvall AH627 rotor at 26 k rpm for 21 hrs at 15°C.

The gradient was then slowly fractioned from the bottom by collecting 15 drop fractions (90

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fractions total). 20 μ l of each fraction was then run on a 1% agarose gel as described above. Fractions containing DNA fragments smaller than 18 kbp and larger than 15 kbp were pooled (fractions 32-40) and dialyzed against TE (10 mM Tris.HCl pH 7.5, 1 mM EDTA pH 8.0) to remove the sucrose. The DNA (3.5ml) was then precipitated with ethanol and resuspended to about 15 μ l (1 mg/ml) under vacuum and stored at -20°C.

EcoRI Arms of bacteriophage lambda-Dash were obtained from Vector Cloning Systems (StrataGene) and were ligated at a concentration of 200 μ g/ml to Mycoplasma target DNA at a concentration of 25 μ g/ml in a total volume of 10 μ l using T4 ligase (Boehringer GmbH) at a concentration of 100 units/ml. The ligation reaction was incubated at room temperature for 2 hours. 4 μ l of the ligation was then packaged into lambda particles using the in vitro packaging kit Gigapack (StrataGene). The phage was then titered on E. coli strain P2392 (StrataGene) and found to be 7.75 x 10^5 pfu/ml (3.1 x 10^5 pfu/ug of lambda-Dash).

E. Screening of Library

The library is screened using the plasmid previously obtained which shows greater than 50% homology to pEG22, by the previously described probing procedure. DNA from positive recombinants is prepared, digested with EcoRI, analyzed by gel electrophoresis, to indicate portions of the M. hyopneumoniae genome composed of several EcoRI restriction fragments. One of the fragments is digested with EcoRI, ligated to EcoRI digested pWHA148 and transformed into E. coli strain JM83 and called pMYCO16; its DNA was prepared and digested with a number of different restriction endonucleases in order to derive the restriction map shown in Figure 3.

Plasmid pWHA148 is prepared by inserting a synthetic oligonucleotide into the Hind III site of pUC18. The amino terminal coding sequence of the X-complementing peptide of B-galactosidase is shown in Figure 4, and contains 8 additional restriction sites over the parent pUC18. The oligonucleotide insert into pUC18 is shown in Figure 4 between the Sph1 and Hind III sites.

An N-terminal portion of pEG22 is used by Southern analysis to hybridize to the 0.6kb AccI-AsuII restriction fragment of pMYCO16. DNA sequence analysis of the 0.6 kb fragment identifies that start codon of the homologous gene.

On the restriction map of pMYCO16 (Figure 3
the gene begins within the 0.6 kb AccI-AsuII
restriction fragment, extends clockwise within the 0.4
kb AsuII - ClaI, 1.2 kb ClaI - ClaI, and 1.4 kb ClaIHindIII fragments, and ends short of the HindIII site.
DNA sequence analysis shows that pMYCO16 contains a
74.5 kD protein homologous to the 70 kD T. cruzi heat
shock antigen.

The DNA-amino acid sequence of the 74.5 kD gene is shown in Figure 5.

F. Expression of full length M.hyo. 74.5 kD antigen in E. Coli

Plasmid pMYC016 DNA (Figure 3) was digested with AccI, treated with Mung Bean nuclease to remove the single stranded AccI tails, re-ligated to delete the 1.9 kb AccI fragment in front of the 74.5 kD antigen gene and transformed into E. coli strain JM83. One transformant was named pMYC029; its DNA was digested with a number of different restriction endonucleases in order to derive the restriction map shown in Figure 6.

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pMYC029 was subjected to DNA sequence analysis which showed that a spontaneous deletion had occured at the ligation juncture, where two bases were deleted and the PstI site was retained, as shown below (only a portion of the 5' to 3' strands are represented).

pMYCO29 expected: TTGCATGCCTGCAGGTACTTTCTTTTGTCT

PstI

pMYCO29 observed: TTGCATGCCTGCAGGCTTTCTTTTGTCT

10 PstI

This fortuitous deletion allows the in frame insertion into the pUC9 open reading frame. Plasmid pMYCO29 is a low level expression plasmid.

G. Construction of pMYCO31 and expression of 74.5 kD antigen fragment

Because the mycoplasma insert of pMYCO29 is oriented away from the Lac promoter of pWHA148, it was desired to insert the gene into another expression vector, pUC9. The two base deletion enabled the gene for the 74.5 kD antigen to be placed in the same reading frame as the beta-galactosidase gene of E. coli vector pUC9.

In order to perform this construction, pMYCO29 DNA was digested with PstI and EcoRI, the PstI - EcoRI fragment containing the entire 74.5 kD coding sequence was purified, ligated to the PstI and EcoRI digested vector pUC9, and transformed into E. coli strain JM83. One transformant was named pMYCO31 (Figure 7); its DNA was prepared and transformed into E. coli strain W3110 by the transformation procedure described above.

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H. Construction of pMYC032

It is known that TGA codons encode the amino acid tryptophan in mycoplasma but normally terminate peptide chain elongation in E. coli and that the trpT176 gene, a mutant tryptophan t-RNA which recognizes UGA (Raftery, et al., Jour. Bacteriol., 158:849-859), allows peptide chain elongation at TGA codons in E. coli laboratory mutants. We reasoned that the addition of trpT176 to expression vectors would allow E. coli peptide chain elongation at the mycoplasma TGA codons of cloned genes.

Plasmid pCAM101 was purchased from James Curran (University of Colorado) as a convenient source of the trpT176 gene and is shown in Figure 8.

DNA from pCAM101 was digested with EcoRI, the
0.3 kb EcoRI fragment which contains the trpT176 gene
was purified, ligated to EcoRI digested pMYCO31, and
transformed into E. coli strain W3110. One
transformant was named pMYCO32 and its restriction
ap is shown in Figure 9.

I. Expression of M. hyopneumoniae 74.5 kD antigen in E. coli

A W3110 (pMYCO32) transformant was selected, grown in L-broth, lysated as previously described, and a portion subjected to polyacrylamide gel electrophoresis. New 75 kD and 43 kD proteins were identified by gel electrophoresis which represented approximately 5% and 0.1% of total E. coli protein, respectively. The pMYCO32 75 kD protein was shown by Western blot to react with the previously described pig antisera raised against the 74.5 kD M. hyopneumoniae antigen.

An improved expression plasmid pMYCO87 has been deposited with the ATCC on June 30, 1989 as ATCC number 68030. It contains an in vitro change of TGA to TGG (Tryptophane) at codon position 211 (see Figure 5).

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J. Use of the recombinant form of Mycoplasma hyopneumoniae 74.5 kD antigen as a vaccine

A W3110 (pMYCO32) transformant from Example 2 was selected, grown in M-9 minimal medium in a 14 liter Chemap fermenter to a cell density of 110 O.D. 600, and 120 g (wet weight) of cells were harvested from 500 ml by centrifugation. A suspension was prepared consisting of 2.3 g of cells per 10 ml of PBS containing 12 mM EDTA, 0.5 mg/ml lysozyme. suspension was incubated at 25 °C for 15 minutes. sonicated on ice for 2 minutes in 30 second bursts, centrifuged at 13,000 g for 10 minutes at 4°C, and the soluble fraction reserved as product. A portion of the product was subjected to polyacrylamide gel electrophoresis. The recombinant form of 74.5 kD antigen made up approximately 25% of the soluble protein and the yield dosages were prepared in PBS at 100 and 500 μ g per dose and emulsified on ice with equal volumes of Freund's incomplete adjuvant (Sigma) immediately prior to use.

Vaccination Test

- Week 0 Three litters of Hampshire, Hampshire X
 Duroc, and York piglets taken by Caesarian
 section.
- 25 Week 1 Piglets divided randomly into 7 pig dosage groups and each vaccinated sub-cutaneously in leg.
 - Week 3 Booster vaccination, as above, opposite leg.
- 30 Week 8 Challenge administered by trans-tracheal inoculation of 106 CCU Mycoplasma hyopneumoniae.
 - Week 12 Necropsy of experimental animals and infection controls.

The	results	were	as	follows:

	Group	Incidence*	Severity**
	Control	5/5	12.4 ± 4.7
	100 ug 74.5 kD	1/4	4.2 <u>+</u> 4.9
5	100 ug recomb. 74.5 kD	2/6	9.7 ± 11.7
	500 ug recomb. 74.5 kD	4/4	25.0 ± 6.1

^{*} Number of pigs with a lung lesion score greater than 5%

10 ** % of lung surface effected (mean \pm std. dev.)

Example 3. -- The 70 kD Hsp Analog from Mycoplasma Gallisepticum.

A. Preparation of Genomic Libraries

Two strains of M. gallisepticum F-K810 and R,

were obtained from R. Yamamoto (U. C. Davis) and grown
in F-80 media for the preparation of genomic DNA. (Nord
Veterinaermed. 27:337-339).

Approximately 22 ml of stationary phase M.
gallisepticum culture was centrifuged at 13,000 X g at
4°C for 10 minutes to harvest mycoplasma cells. The
supernatant was discarded and the cell pellet was
resuspended in PBS to wash. Cells were harvested by
centrifugation after washing. The cells were washed a
total of three times with PBS and the resulting cell
pellet frozen at -78°C. After thawing, the cells were

- perfet frozen at -78°C. After thawing, the cells were resuspended in 2 ml 10 mM Tris-HCl pH 8.0, 50 mM EDTA, 1% SDS, and 100 μ g Proteinase K was added. The cells were lysed at 50°C for one hour with occasional mixing. The lysate was extracted with phenol then with
- chloroform/isoamyl alcohol to remove cellular debris. The DNA-containing aqueous phase was dialyzed against 4 liters of 10 mM Tris-HCl, 5 mM EDTA twice, and 10 mM Tris-HCl, 1 mM EDTA once. From each strain, 60 μg of DNA was recovered, an amount sufficient for restriction
- analyses. Southern blot analyses, and library construction. Restriction digests indicated that the

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two strains are similar to each other with limited restriction fragment length polymorphism.

B. <u>Mixed oligonucleotide probes for isolating the Hsp70 protein from M. gallisepticum</u>

When the Hsp70 amino acid sequence from T. Cruzi aligned with the amino acid sequence of the M. hyopneumoniae 74.5 kD antigen of Example 2. Several regions containing sequences six amino acids in length are identical in both sequences. The array of DNA sequences which could encode these amino acid regions was determined. The two amino acid sequences corresponding to nucleotide sequences having the lowest degeneracy, were selected for use as oligonucleotide probes. These were synthesized as follows:

15 COD1159 Ile-Ile-Asn-Glu-Pro-Thr ATA-ATA-AAC-GAA-CCA-AC C C T G C

T T G

20 COD1218 Gly-Gly-Gly-Thr-Phe-Asp GGA-GGA-GGA-ACA-TTC-GA

> CCCCT GGGG TTTT

Pools of the above oligonucleotides were labeled with ³²P using polynucleotide kinase (BRL) and used to probe Southern transfers of HindIII digested M. gallisepticum chromosomal DNA. After 50°C washes in 6X NET, 0.1 SDS, COD 1159 hybridized to two HindIII fragments. COD 1218 hybridized to two HindIII fragments at 45°C under likewise identical conditions. Both probes hybridize to an apparently identical 3.4 kb fragment, where as the other fragments differ in length

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and probably represent hybridization due to non-specific sequence homology. The hybridization of both probes to the same 3.4 kb HindIII fragment is highly significant as the probability that hybridization of both probes to the same fragment of genomic DNA results from non-specific sequence homology is less that 2X10⁻³. The hybridization patterns for DNA purified from strain R strain and F-K810 strain of M. gallisepticum were identical to one another.

Plasmid DNA from pMYCO87, containing the gene for M. hyopneumoniae (ATCC 68030 deposited with the American Type Culture Collection on June 30, 1989) was labeled using the Boeringer Mannheim nonradioactive Southern hybridization kit (Genius kit) and used to probe a Southern transfer of EcoRI and HindIII restriction digested chromosomal DNA from the F-strain and M. hyopneumoniae as a positive control. The probe detected bands of the expected size in the M. hyopneumoniae genome and an EcoRI band of 6.8 kb and a Hind III band of 3.3kb in the M. gallisepticum digests after washes at 65°C in 0.5X SSC and 0.1% SDS.

C. Preparation of Size Selected Genomic Libraries

The general approach for cloning the hsp antigen gene from M. gallisepticum was analogous to the procedure used for the T. cruzi 70 kD hsp. M. gallisepticum genomic DNA, 1 µg from both the R strain and the F-K8 I O strain, was digested to completion with the bacterial restriction endonuclease HindIII and separated on 3.25% polyacrylamide gels. DNA from four gel slices containing restriction digest fragments between 2 and 5 kb was electroeluted. An aliquot of DNA electroeluted from each of the four gel slices was subjected to agarose gel electrophoresis, transfered to a nitrocellulose membrane by Southern transfer and probed with ³²P-labeled COD1159 to identify the fraction which contains the 3.3kb hybridizing HindIII band. In

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this way, a positive DNA fraction was identified. This positive DNA fraction was then ligated into Hind III digested pUC9 and transformed into E. coli DH5a.

D. <u>Identification of Positive Clones</u>

For each strain, 12 and F-K810, plasmid DNA from forty-eight recombinant clones was isolated by the method of Holms and Quigley 1981 (Anal. Biochem. 114:193-197, 1981), transferred to nitrocellulose using a Bio-Rad dot blot apparatus, and probed with COD1159 in the case of the R-strain or both COD1159 and COD1218 on duplicate blots In the case of strain F-K810.

DNA from pCAM101 was digested with EcoRI, a 0.3 kb EcoRI fragment including trpT176 was purified, ligated to EcoRI digested pUC9, transformed into E. coli strain JM83, and one transformant was named pWHA160 (see Figure 12).

Plasmid pMGA4 DNA was digested with HindIII and BgIII, ligated to HindIII and BamHI digested pWHA160, digested with BamHI and BgIII, and transformed into E. coli strain DH5a. One transformant was named pMGA10. The MGA10 transformant was grown in L-broth at 37°C, and the cells harvested by centrifugation and frozen. The cell pellet from 4 ml of culture was resuspended in 100 μ l of a solution consisting of 0.5 mg/ml hen egg-white lysozyme dissolved in 25 mM Tris pH 8.0 10 mM EDTA; and incubated at 25°C for 10 minutes.

A portion of the resulting lysate was subjected to polyacrylamide gel electrophoresis and a new 67 kD protein was identified. Western blot analysis, using pig anti-74.5kD serum, showed that the new 67 kD protein was immunologically related to Hsp70.

F. <u>Use of Bacterially Produced M. gallisepticum Hsp 70</u> Protein to Raise an Immune Response in Chicken

The purified M. gallisepticum protein is concentrated by lyophilization and resuspended to a final concentration of 0.5-2.0 mg/ml in 0.1% SDS. For use, the immunizing antigen is formulated in one volume of protein concentrate to three volumes of oil carrier consisting of 5% Arlacel, 94% Drakeol 6-VR and 1% Tween 80. The dose of the antigen employed is 100 µg/dose. Chicken receive the formulated vaccine by subcutaneous injection. A booster vaccination by the same route is

Numerous modifications and variations of the present invention are possible in light of the above teachings; therefore, within the scope of the appended claims the invention may be practiced otherwise than as particularly described.

done two weeks later.

Table 1. Representative Pathogenic Organisms.

1: DISEASE AGENTS 1.1: BACTERIA 1.1.1: ACTINOBACILLUS SPP. 1.1.1.1: Actinobacillus lingiresii Mastitis infections in cattle, swine, equine 1.1.1.2: Also known as Haemophilus swine pneumonia 1.1.2: BACILLUS SPP. Bacillus anthracis Anthrax, an acute febrile disease of all mammals 1.1.3: BORDETELLA SPP. B. bronchiseptica - repiratory disease in 1.1.3.1: many species B. pertussis - whooping cough in man 1.1.3.2: 1.1.4: BORRELIA SPP. B. burgdorferi - Lyme disease in dogs, 1.1.4.1: deer, man 1.1.5: BRUCELLA SPP. Brucella abortus, B. suis, B. melitensis 1.1.5.1: brucellosis in cattle, sheep, swine, equine, canine, man 1.1.6: CAMPYLOBACTER SPP. Campylobacter fetus 1.1.6.1: infertility and embryonic causes death in cattle, swine, sheep, equine (vibriosis) Vibrio cholerae - cholera in man 1.1.6.2: 1.1.7: CHLAMYDIA SPP. C. psittaci - respiratory disease in 1.1.7.1: birds C. cati - conjunctivitis in cats 1.1.7.2: 1.1.8: CLOSTRIDIUM SPP. C. chauvoei 1.1.8.1.: blackleg in cattle and sheep 1.1.8.2: C. septicum malignant edema in cattle and sheep C. haemolyticum 1.1.8.3: red water in cattle C. novyi 1.1.8.4: black disease in cattle and sheep C. sordelli 1.1.8.5: big head disease in cattle and sheep C. perfringens 1.1.8.6: enterotoxemia in cattle, sheep, swine,

equine, gas gangrene in man

3.

1.1.8.7: C. tetani tetanus in all mammals 1.1.8.8: C. boutulinum 8 types, causing botulism in all species 1.1.9: CORYNEBACTERIUM SPP. 1.1.9.1: C. diptheria - Diptheria in man C. pyogenes -causes pyogenic processes in 1.1.9.2: cattle, sheep, swine, goats C. renale - cystitis in cattle 1.1.9.3: 1.1.9.4: C. equi - pneumonia in horses 1.1.10.1: ERYSIPELOTHRIX SPP. Erysipelothrix rhusipothiae - erysipelas 1.1.10.1: in swine and man 1.1.11: HAEMOPHILUS SPP. H. influenza, respiratory disease 1.1.11.1: various species H. paraninfluenza, H. parasuis, H. suis -1.1.11.2: respiratory disease in swine 1.1.12: KLEBSIELLA SPP. 1.1.12.1: Klebsiella pneumoniae - Pneumonia and septicemia in animals and man 1.1.13: LISTERIA SPP. 1.1.13.1: Listeriosis monocytogenes encephalitis in ruminants 1.1.14: MYCOBACTERIUM SPP. M. tuberculosis, M. bovis, M. avium -1.1.14.1: Tuberculosis in various species M. paratuberculosis - Johne's disease in 1.1.14.2: cattle, sheep, and goats 1.1.15: PASTEURELLA SPP. P. pestis - Plague in man and rodents 1.1.15.1: multocida haemolytica, P. 1.1.15.2: respiratory disease in many species 1.1.16: PSEUDOMONAS SPP. P. aeruginosa - respiratory disease in 1.1.15.1: various animals P. mallei - Glanders disease in dogs and 1.1.16.2:

S. typhimurium - enteric disease in a

S. typhisuis, S. choleraesuis - enteric

cats

number of species

disease in swine

S. typhi - Typhoid fever

1.1.17: SALMONELLA SPP.

1.1.17.1:

1.1.17.2:

1.1.17.3:

1.1.17.4: 1.1.17.5: 1.1.17.6:	S. paratyphi - Paratyphoid - A in man S. gallinarum - fowl typhoid S. pullorum - pullorum disease in chickens
1.1.18: STRE 1.1.18.1: 1.1.18.2: 1.1.18.3: 1.1.18.4: 1.1.18.5:	SPTOCOCCUS SPP. S. agalactiae, S. dysgalactiae - mastitis in numerous species S. dispar - enteritis in numerous species S. equi - cholic in horses S. genitalium - uterine infections in horses S. pneumoniae - respiratory disease in man
1.1.19.1: 1.1.19.2:	PHYLOCCUS SPP. S. aureus - mastitis in many species S. epidermidis - pyoderma in many species
1.1.20: TULA 1.1.20.1:	AREMIA SPP. Francisella tularensis - Tularemia in man
1.2.6: HERPE 1.2.6.1: 1.2.6.2: 1.2.6.3: 1.2.6.4: 1.2.6.5: 1.2.6.7: 1.2.6.8: 1.2.6.9: 1.2.6.10: 1.2.6.11:	ESVIRIDAE H. simplex Type 1 - Oral Herpes in man H. simplex Type 2 - Genital Herpes in man Epstein-Barr Virus - Mononucleosis in man H. smiae - Herpes B. in primates H. suis-Adjuskie's disease - pseudorabies in swine and cattle H. canis - Respiratory infection of dogs H. equi - Equine rhinopneumonitis - respiratory and abortion in horses H. bovis - IBR (Infectious Bovine Rhinotracheitis) in cattle H. felis - FVR (Feline Viral Rhinotracheitis) Laryngotracheitis virus - Laryngotracheitis virus - Laryngotrachetis in birds Marek's Disease Virus - Merek's disease in birds Feline calicivirus (FCV) Cytomegaloviruses-many diseases in various animals
1.2.13: POX 1.2.13.1: 1.2.13.2: 1.2.13.3: 1.2.13.4: 1.2.13.5: 1.2.13.6:	SMALLPOX - WAS A MAJOR DISEASE IN MAIN VACCINIA - USED TO VACCINATE AGAINST SMALLPOX

1.2.13.7:	CARDIBOVIUM
1.2.20.7.	CAPRIPOXIVIRUSES - LUMPY SKIN DISEASE IN SHEEP AND GOATS
1:2:13:8:	PARAPOXIVIRUSES - "SORE MOUTH" IN SHEEP
	AND GOATS, BOVINE PAPULAR STOMATITIS
*	
1.3: MYCOPLAS	
1.3.1:	M. mycoides - Bovine respiratory disease
1.3.2: 1.3.3:	M. bovis - bovine mastitis
1.3.4:	M. bovigenitalium - bovine epidymitis
1.0.1.	M. bovoculi - Infectious bovine keratoconjuntivitis
1.3.5:	M. bovirhinis and M. dispar - respiratory
	disease
1.3.6:	M. hyorhinis and M. hyosynoviae -
	respiratory disease and lameness in swine
1.3.7:	m. gallisepticum and M. synoviae -
	respiratory disease in poultry
1.4: RICKETTS	STA
1.4.1:	Rickettsiaceae
1.4.1.1:	R. prowszekii - Typhus fever
1.4.1.2:	R. typhi - murine thyphus in man
1.4.1.3:	R. rickettsii - Rocky Mountain Spotted
	Fever
1.4.1.4:	Coxiella Burnetii - Q Fever in cattle.
	sheep, goats, birds, and man
1.4.1.5:	Cowdria ruminatum - Heartwater in cattle
1.4.2: 1.4.2.1:	Anaplasmataceae
1.7.2.1:	A. marginale and A. centrale - Anaplasmosis in cattle
1.4.2.2:	A. ovis - Anaplasmosis in sheep
1.4.2.3:	Haemobartonella felis - Hemobartonellosis
	in cats (Feline Infectious Anemia)
1.4.2.4:	Haemobartonella canis - Hemobartonellosis
	in dogs
1.4.2.5:	Eperythrozoon - parasites which attack
	red blood cells in various animals
1.5: CHLAMYDI	t some of
1.5.1:	
1.3.1:	pulonary disease in man and birds
1.5.1.1:	also causes Sporadic Bovine
	Encephalomyelitis and polyarthritis in
	cattle
1.5.1.2:	also causes Epizootic Abortion in cattle
	and sheep
1.5.1.3:	also causes pneumonia in cattle and hseep
1.5.1.4:	also causes Feline Pneumonitis in cats
1.5.2:	C. trachomatis - Veneral disease in man
1.6: SPIROCHA	AFTAT.F
1.6.1:	Leptospria spp.

1.6.1.1:	L. canicola, L. grippotyphosa, L. hardjo,
1.0.1.1.	7
1.6.1.2:	L. pomona - all cause disease in various
1.0.1.2.	species
1.6.2:	Transpars SPP.
	T. hyodysenteriae - Swine Dysentery
1.6.2.1:	T. pallidum - Syphilis in man
1.6.2.2:	Borrelia spp.
1.6.3:	B. anserina O Avian borrelosis or
1.6.3.1:	B. anserina o Avian Dollous
•	spirochaetosis in birds
1.7: FUNGAL I	DISEASES Asperigillus fumigatus - brooder
1.7.1:	Asperigillus rumigatus
1.7.2:	Blastomyces dermatitidis - pulmonary
	infection in animals and man
1.7.3:	Candida albicans - Thrush in birds, cats,
	cattle, swine and man
1 7 4. EPIDE	BMADUVTAN SPP
1.7.4.1:	E. floccosum - Athletes foot in man
1.7.5: HISTO	STACMA CDD
1.7.5.1:	H. capsulatum - systemic fungal infection
1.7.5.1:	in many species
1.7.6: MICRO	CDADIM SPP
1.7.6: MICRO	M. canis - ringworm in dogs, cats, man,
1.7.6.1:	
	M. gypseum - ringworm in dogs, cats,
1.7.6.2:	M. gypseum - Illigworm
	horses, man
1.7.7: TRICH	OPHYTON SPP. T. rubrum - ringworm in dogs, primates,
1.7.7.1:	T. Pubrum - Pingwolm in dogs, Polimers
1.7.7.2:	and man T. equinum and T. quinkeanum - ringworm
	in horses
1.7.8: MYCO	TOXICOSES (Moldy feed) caused by numerous
filamentous	finati
1.7.8.1:	Aflatoxins, Mycotoxins, Aspergillus
1.7.0.2.	toxins
2: PARASITES	
Z: PARASITE	, 11
2.1: PROTOZO	/n A
2.1.1: AMEB/	Entamoeba histolytica - Amebic dysentery
2.1.1.1:	in dogs, cats, pigs and man
2.1.2: BABES	Babesia bigemina and B. bovis are major
2.1.2.1:	L_L_AAAA4
	(babesiosis also known as Texas fever,
•	(babes10515 also Allowit as Island
	Tick Feber, Prioplasmosis)
2.1.2.2:	B. argentina, B. Divergens, and B. major
~· • · ~ · ~ ·	also cause babesiosis in cattle
2.1.2.3:	R canis and B. Gigsoni - Cause
4.1.2.3.	babesiosis in dogs
	

2.1.2.4:	B. equi and B. caballi cause babesiosis
	in horses
2.1.2.5:	B. motasi and B. ovis - cause babesiosis
	in horses
2.1.2.6:	B. trautmanni - babesiosis in pigs
2.1.2.7:	B. felis - babesiosis in cats
2.1.3: COCCI	
2.1.3.1:	EIMERIA SPP.
•	E. tenelia, E. necatrix, E. brunetti, E. acervulina, E. maxima in chickens
	acervulina, E. maxima in chickens
	E. bovis, E. zuernii in cattle
2.1.3.2:	ISOSPORA SPP.
	I. suis - seine
2.1.3.3:	SARCOYSTIS SPP.
	S. tenella - infects sheep
	S. blanchardi, S. fayerei, and S.
	fusiformis - infect cattle
	S. miescheriana - infects swine
2.1.3.4:	TOXOPLASMA GONDII
	wide spread distribution, especially in
	cats, swine, sheep, humans
	causes abortion, birth defects, deafness
2.1.3.5:	CRYTOSPORIDUM SPP.
2.1.3.3.	cause diarrhea in cattle, swine, sheep,
	birds, and man
	A component of AIDS complex
2.1.4: GIARI	
	G. lamblia - infects man
2.1.4.1:	G. canis - infects dogs
	G. cati - infects catas
	G. bovis - infects cattle
2.1.5:	LEISHMANIA SPP.
2.1.5.1:	L. donovani - visceral leishmania in man,
	dogs, cats, cattle sheep
2.1.5.2:	L. tropica - cutaneous leshmania in man,
	dogs, and rodents
2.1.5.3:	L. braziliensis - American leishmaniasis
	in man, dogs, and cats
2.1.6: PLASE	MODIUM SPP.
2.1.6.1:	Plasmodium falciparum - malaria in man
2.1.6.2:	P. malariae, P. vivax, and P. ovale -
	malaria in man
2.1.6.3:	P. gallinaceum - avian malaria
2.1.6.4:	numerous Plasmodium spp. cause malaria in
	man
2.1.7: PNEU	MOCYSTOSIS SPP.
2.1.7.1:	P. carinii - cause of pneumonia in man,
	dogs, horses, swine, goats
2.1.7.2:	A component of the AIDS complex
2.1.7.2: 2.1.8: THEI	
	T. parva, T. annulata, T. mutans, T.
2.1.8.1:	lawrencei and T. cervi
	TOMICINGS WING TO ACTAT

all cause East Coast Fever in cattle. buffalo and deer 2.1.8.2: T. hirci and T. ovis infect sheep 2.1.9: TRITRICHOMONAS SPP. 2.1.9.1: T. vaginalis - a veneral disease of man 2.1.9.2: foetus - causes trichomonaiasis, genital infection of cattle 2.1.9.3: Trichomonas gallinae tricomoniasis, a G.I. infection in birds 2.1.10: TRYPANOSOMA SPP. 2.1.10.1: T. cruzi - Chagas disease in man 2.1.10.2: T. congolense -- Trypanosomiasis cattle, horses, pigs, dogs gambiense 2.1.10.3: rhodesiense and T. sleeping sickness in man and antelope 2.2: HELMINTHS 2.2.1: TREMATODES 2.2.1.1: FLUKES Fasciola hepatica - cattle and sheep F. gigantica - cattle and sheep Fascioloides magna - cattle, sheep and swine Dicrocoelium dendriticum - cattle, sheep, horses, swine, man SCHISTOSOMIASIS 2.2.1.2: Schistosoma japonicum, S. hematobium, S. mansoni, S. intercalatum - man S. bovis, S. spindale, S. mattheei - cattle, sheep, goat, horse S. nasalis, S. indium - cattle, sheep, goats PARAGONIMIASIS (SALMON POISONING) 2.2.1.3: Paragonimus westermani - man P. kellicotti - mink, dog, cat, pig 2.2.2: CESTODES 2.2.2.1: **TAPEWORMS** Taenia saginata, and T. solium man (cysticercus) granulosus. Echinococcus and E. multilocularis - man, dog Taenia hydatigena, T. ovis - dog T. pisiformis - dog and cat Dipylidium caninum - dog and cat Anoplocephala magna, A. perfoliata horses ECHINOCCUS SPP. 2.2.2.2: DIPHYLLOBOTHRIUM SPP. 2.2.2.3: SPIROMETRA SPP. 2.2.2.4: FASCIOLA SPP. 2.2.2.5: 2.2.3: NEMATODES FILARIAL PARASITES 2.2.3.1: Dirofilaria immitis - heartworm in dogs HOOKWORMS 2.2.3.2:

	A. duodenale and Necator americanus -
	hookworm in man
	A. caninum, A. braziliense - dogs and
	cats .
	Uncinaria stenocephala - dogs
•	Bunostomum phlebotomum - cattle
	B. trigonocephalum - sheep and goats
	Globecephalus urosubulatus - swine
2.2.3.3:	KIDNEY WORMS
•	Dicoctophyma renale - dog
2.2.3.4:	LUNGWORMS
	Dictyocaulus viviparus - lungworm in
	cattle
	D. filaria - lungworm in sheep, goat,
	cattle Muellerium capillaris - lungworm in sheep
	Metastrongylus apri, M. pudendotectus, M.
	salmi - swine
2.2.3.5:	NODULAR WORMS
2.2.3.3:	Oesophagostomum denatum - swine
	O. radiatium, and O. columbianum -
	cattle, sheep, goats
2.2.3.5:	ONCHOCERIASIS
2.2.0.0	Onchocerca volvulus - blindness in humans
2.2.3.7:	PINWORMS
	Enterobius vermicularis - man
	Oxyuris equi - horses
	Skrjabinema ovis - sheep and goats
2.2.3.8:	ROUNDWORMS
	Ascaris lumricoides - roundworms in man,
	swine
	Toxocara canis - dogs Toxocara cati - cats
	Parascaris equorum - horse
	Ascaridia galli - chickens
2.2.3.9:	SPIROCERCAS
4.2.3.9:	Spriocerca lupi - dogs
2.2.3.10:	STOMACH WORMS
2.2.3.20.	Habronema, H. majus, H. megastoma -
• •	horses
2.2.3.11:	STRONGYLES
	Strongylus vulgaris, S. equinus, S.
	edentatus - horses
2.2.3.12:	STRONGYLOIDS
	Strongyloides westeri - horses
	S. stercoralis - man
	S. ransomi - swine
	S. canis - dogs S. tumefaciens - cats
2.2.3.13:	TRICHINA Trichinella spiralis - trichinella in
	swine and man
2 2 2 34.	TRICHOSTRONGYLES
2.2.3.14:	

Ostertagia ostertagi - cattle Haemonchus placei - cattle Trichostronglyus axei - cattle Cooperia punctata - cattle Haemonchus contortus, Cuperia curticei sheep Ostertagia circumcincta - sheep Trichostronglyus colubriformis - equine, swine, cattle, sheep Nematodirus filicollis - cattle and sheep Hyostrongylus rubidus - swine WHIPWORMS 2.2.3.15: Trichuris ovis - cattle, sheep, goats Trichuris suis - swine T.. trichiura - man T. vulpis - dogs ARTHROPODS 2.3: ACARIASIS 2.3.1: Demodex folliculorum - mange in dogs, 2.3.1.1: cats, cattle, swine, sheep, man Demodex phylloides - mange in swine 2.3.1.2: Dermacentor andersoni - wood tick Dermanyssus gallinae - red m 2.3.1.3: mite 2.3.1.4: poultry Ixodes holocyclus - Austrailian tick 2.3.1.5: Notoedres cati - cat mange 2.3.1.6: Otobius megnini - spinose ear tick 2.3.1.7: Ostodectes cynotis - ear mite in dog, cat 2.3.1.8: Psoroptes communis - scab in cattle, 2.3.1.9: sheep, horses Sarcoptes scabiei, S. canis - mange in 2.3.1.10: dogs DIPTERA 2.3.2: BOTFLIES 2.3.2.1: equine intestinalis Gasterophilus botfly hemorrhoidalis -Gasterophilus nose botfly Gasterophilus nasalis - equine chinfly Gasterophilus pecorum - European botfly Gasterophilus inermis - botfly Oestrus ovis - sheep botfly FLEAS 2.3.2.2: Otenocephalides canis - dog flea Ctenocephalides felis - cat flea FLIES 2.3.2.3: Chrysops spp. - deer flies Fannia spp. - little house flies Haematobia irritans - horn flies Haematotobia irritans exigua - buffalo fly (similar to horn fly) Hermetia illucens - black soldier fly Hybomitra spp. common fly

Hydrotaea irritans - head flies Ophyra spp. - dump flies Melophagus ovinus - sheep ked Musca autumnalis - face flies Musca domestica - house fly Muscina spp. - false stable flies Simulium spp. - black flies (no-see-ums) Stomoxys calcitrans - stable flies Tabanus spp. - horse files 2.3.2.4: **GRUBS** Hypoderma lineatum, H. bovis - Heel fly, cattle grub Calitroga americana - screw-worm fly Dermatobia hominis - cutaneous myiasis in man, cattle sheep, dogs, cats Cochliomyia hominivorax - blow fly 2.3.2.5: LICE Damalinia bovis - cattle biting louse Anoplura spp. - cattle louse Haematopinus eurysternus shortnosed cattle louse Linognathus vituli - longnosed cattle louse Solenoptes capillatus little blue cattle louse Haematopinus suis - swine lice Haematopinus asini - horse sucking louse Trichodectes canis - dog louse Felicola subrostrata - cat louse MOSQUITOES 2.3.2.6: Aedes spp. Anopheles spp. Culex spp. Culiseta spp. Psorophora spp.

Disease Pathogen(s)

Malaria Plasmodium falciparum

P. vivax
P. malariae
P. ovale
P. berghei
etc.

Chagas' Disease

Trypanosoma cruzi

African Trypanosomiasis Trypanosoma gambiense

T. rhodesiense

T. brucei

etc.

Leishmaniasis

Leishmania donovani

L. infantum
L. tropica
L. mexicana
L. braziliensis
L. chagasi
etc.

Leprosy Mycobacterium leprae

Tuberculosis Mycobacterium tuberculosis

Filariasis Brugia malayi

B. timori

Onchocerca volvulus Wuchereria bancrofti

Schistosomiasis Schistosoma mansoni

S. japonicum

Leptospirosis Leptospira interrogans

L. iceterohaemorrhagiae

L. hebdomadis L. pomona

etc.

Plaque Yersinia pestis

Typhoid Fever Salmonella typi

Cholera Vibrio cholerae

Diptheria Corynebacterium diphtheriae

Lyme Disease Borrelia burgdorferi

Pneumonia/bronchitis Streptococcus pneumoniae

Mycoplasma pneumoniae Branhamella catarrhalis Bordetella bronchiseptica

Haemophilus influenza

Urethritis Mycoplasma hominis

Ureasplama urealyticum

Giardia Giardia lamblia

Amoebic dynsentery Entamoeba histolytica

Syphilis Treponema pallidum

Chlamydia Chlamydia trachomatis

Candidiasis Candida albicans

C. glabrata

Gonorrhea Neisseria gonorrhoeae

Toxoplasmosis Toxoplasma gondii

Tetanus Clostridium tetani

Caries Streptococcus mutans

Whooping cough Bordetella pertussis

Q fever endocarditis Coxiella burnetti

Anthrax Bacillus anthracis

Brucellosis Brucella abortus

Numerous modifications and variations of the present invention are possible in light of the above teachings; therefore, within the scope of the appended claims the invention may be practiced otherwise than as particularly described.

WHAT IS CLAIMED IS:

- 1. A vaccine for protecting against an organism, comprising:
- (a) a protein which is capable of eliciting an antibody which recognizes at least one epitope of a native protein present in the organism, said native protein having at least 50% homology with a heat shock protein of T. cruzi; and
 - (b) a physiologically acceptable carrier.
- 2. A vaccine of claim 1 wherein the native protein is derived from a species of Mycoplasma, Mycobacteria, or Trypanosoma, provided that the native protein is not derived from Trypanosoma cruzi.
- 3. A vaccine of claim 2 wherein the native protein is derived from a species of Mycoplasma selected from the group consisting of M. mycoides, M. bovis, M. bovigenitalium, M. bovoculi, M. bovirhinis, M. dispar, M. hyorhinis, M. hyosynoviae, M. hyopneumoniae, M. gallisepticum, M. pneumoniae and M. synoviae.
 - 4. A vaccine of claim 2 wherein the native protein is derived from a species of Mycobacteria selected from the group consisting of M. bovis, M. leprae and M. tuberculosis.
- 5. A vaccine of claim 3 wherein the native protein is derived from a species of Mycoplasma selected from the group consisting of M. hyopneumoniae and M. gallisepticum.

6. A process for protecting a host against an organism comprising:

administering an effective amount of a protein capable of eliciting an antibody which recognizes at least one epitope of a native protein present in the organism said native protein having at least 50% homology with a T. cruzi heat shock protein.

- 7. A process of claim 6 wherein the native protein is derived from a species of Mycoplasma,

 10 Mycobacteria, or Trypanosoma, provided that the native protein is not derived from Trypanosoma cruzi.
- 8. A process of claim 7 wherein the native protein is derived from a species of Mycoplasma selected from the group consisting of M. mycoides, M. bovis, M. bovigenitalium, M. bovoculi, M. bovirhinis, M. dispar, M. hyorhinis, M. hyosynoviae, M. hyopneumoniae, M. gallisepticum, M. pneumoniae, and M. synoviae.
- 9. A process of claim 7 wherein the native 20 protein is derived from a species of Mycobacteria selected from the group consisting of M. bovis, M. leprae, and M. tuberculosis.
- 10. A process of claim 8 wherein the native protein is derived from a species of Mycoplasma
 25 selected from the group consisting of M. hyopneumoniae and M. gallisepticum.
 - 11. A process for determining an organism in a host comprising:

contacting a sample derived from a host containing or suspected of containing an organism with an antigen which is recognized by an antibody elicited in response to a protein present in the organism, said protein

10

having at least 50% homology with a heat shock protein of T. cruzi; and determining antibody in said sample bound by said antigen.

12. A process for determining an organism in a host, comprising:

contacting a sample derived from a host containing an organism or suspected of containing an organism with an antibody or fragment of said antibody, said antibody recognizing at least one epitope of a native protein present in the host, said native protein having at least 50% homology with a heat shock protein of T: cruzi; and

determining protein present in said organism bound to said antibody.

- 13. A process of claim 12 wherein the native protein is derived from a species of Mycoplasma,
 Mycobacteria, or Trypanosoma, provided that the native protein is not derived from Trypanosoma cruzi.
- protein is derived from a species of Mycoplasma selected from the group consisting of M. mycoides, M. bovis, M. bovigenitalium, M. bovoculi, M. bovirhinis, M. dispar, M. hyorhinis, M. hyosynoviae, M. hyopneumoniae, M. gallisepticum, M. pneumoniae and M. synoviae.
 - 15. A process of claim 13 wherein the native protein is derived from a species of Mycobacteria selected from the group consisting of M. bovis, M. leprae, and M. tuberculosis.

- 16. A process of claim 14 wherein the native protein is derived from a species of Mycoplasma selected from the group consisting of M. hyopneumoniae and M. gallisepticum.
- 5 17. A recombinant sequence of nucleic acid encoding the heat shock proteins of M. hyopneumoniae and M. gallisepticum as depicted in figures 5 and 11, respectively.

FIGURE 1-1

Translation of clone pFP70-47

						aal—			C9 s	eaue	nce-		
								_		_			r leu
	T	ICAC	ACAG	GAAA	CAGC'	YA T	G AC	CAT	G AT	T AC	GCC	A AG	CTTG
9							-			_			>
			-	_									lys
23													AAG
													gly
<i>3</i> 7									GGC				
									gln				
ACA 51	ACT	TAC	TCG	TGC	GIT	GGT	GIT	TGG	CAG	AAC	GAG	CGC	GIG
_					_	_			arg				
GAG 65	ATC	ATT	GCG	AAC	GAC	CAG	GGC	AAC	CGC	ACA	ACG	CCG ·	TCG
_			_		_				leu				
TAC 79	GIG	GCG	TIC	ACC	GAC	ACG	GAG	CGT	CTG	ATC	GGT	GAT	GCC
	_								thr				
GCG 93	AAG	AAC	CAG	GIT	GCG	ATG	AAC	CCG	ACG	AAC	ACC	GIC	TTC
_		_	_				_	_	phe		_		
GAC 107	GCG	AAG	CGC	CTC	ATT	GGG	CGG	AAG	TIC	AGC	GAC	CCC	GTT
													thr
GIG 121	CAG	TCG	GAC	ATG	AAG	CAC	IGG	CCC	TTC	AAG	GTC	ATC	ACG
													gly
AAG 135	GGC	GAC	GAC	AAG	CCG	GIG	ATC	CAG	GTG	CAG	TIC	CGC	GGC
glu	thr	lys	thr	phe	asn	pro	glu	glu	val	ser	ser	met	val
GAG 149	ACA	AAG	ACG	TTC	AAC	CCG	GAG	GAG	GIG	AGC	TCG	ATG	GIG
		_		_	_			_	ser	_			
CIG	TCA	AAG	ATG	AAG	GAG	ATT	GCG	GAG	TCG	TAC	CTG	GGC	AAG



163													
	val	1,70	1170	272	τ <i>τ</i> ລ]	772]	thr	าราลไ	nro	ala	tur	phe	asn
	GIG												
	GIG	DAA	HAG	GCC	GIG	GIG	ACI	GIG		300		110	1410
187		. 7 .		T	-7-	ما ما	3		-1-	~7	+h~	410	272
	ser												
	TCC	CAG	CGG	CAG	حال	ACG	AAG	GAI	GCC	GGC	ACG	AIC	فالم
201		_	_	_					-			- 7 -	- 7 -
	met												
	ATG	GAG	GTG	CIG	CGC	ATC	ATC	AAT	GAG	CCG	ACA	GCT	GCC
215								_	_		_	_	
	ile												
GCC	ATT	GCG	TAC	GGC	CTG	GAC	AAA	GIG	GAG	GAC	GGC	AAG	GAG
239													
arg	asn	val	leu	ile	phe	asp	leu	gly	gly	gly	thr	phe	asp
CGC	TAA	GIG	CTC	ATC	TTT	GAC	CTT	GGC	GGC	GGC	ACG	TTT	GAT
253													
val	thr	leu	leu	thr	ile	asp.	gly	gly	ile	phe	glu	val	lys
	ACG												
267													•
ala	thr	asn	alv	asp	thr	his	leu	qly	gly	glu	asp	phe	asp
	ACG												
281													
_	arg	leu	val	ser	his	phe	thr	asp	alu	phe	lvs	arq	lys
	CGC												
295	000	010	010		— 10								
	lys	വിഗ	1775	asn	len	thr	thr	ser	aln	arq	ala	leu	arq
	AAG												
309		000			010	1100	11021	1200		000	000	010	000
	leu	2 **	+hr	212	CT 7C	~711	2200	212	1770	2777	thr	ا ا ا	gar
_	CIC	_			_	_	_		_	_			
	CIC	CGC	ACC	GU	IGC	CAC	CGC	GCC	DAM	CGC	ACG	CIG	100
323	- 7 -	-1-	1	-1-	_1	47.a	٠.٦.	410		212	7.011	nho.	2000
	ala												
	GCG	GCA	CAG	فالك	ACG	ATT,	GAG	AIC	GAC	فالم	CIG	T.T.T.	GAC
337	_		. 1			<u>. 1</u> .		4-1		_7_		1	7
	val												
AAC	GTG	GAC	TTC	CAG	GCA	ACC	ATC	ACT	CGC	GCC	CGC	TTC	GAG

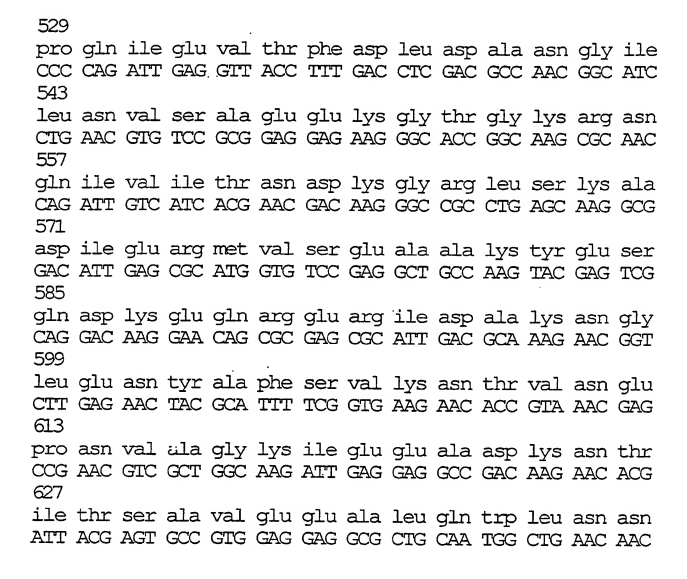
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FIGURE 1-3

351													
glu GAG	leu CTC	cys TGC	GGC GL	asp GAC	leu CTC	phe TTC	arg CGA	GGG Gly	thr ACG	leu CTG	gln CAG	pro CCG	val GIG
365		_		_			-			-			7
	arg CGT												
379	CGI	GIG	CIC	CAG	GAC	GCC	DAG	AIG	CAC	בעער	CO1	000	010
	asp	val	val	leu	val	gly	gly	ser	thr	arg	ile	pro	lys
CAC	GAC	GIG	GTG	CTC	GTC	GGC	GGC	TCC	ACC	CGC	ATT	CCA	AAG
393		_	_	_			,	,	. 7	-	7		1
	met ATG												
407	AIG	CAG	CIG	GIG	101	GAC	777	110	661	GGC	MO	CERT	CIO
	lys	ser	ile	asn	pro	asp	glu	ala	val	ala	tyr	gly	ala
	AAG												
421	_	_	_			-		-	. 7	7		7	
	val GTG												
435	GIG	CAG	GCC	110	AIC	CIG	ACG	990	GGC	בעת	AGC	MO	CAC
	glu	gly	leu	val	leu	leu	asp	val	thr	pro	leu	thr	leu
	ĞAG												
449			_	_	_	_	_				-	• -	-
gly	ile	glu	thr	ala	gly	gly	val	met	thr	ser	Leu	TTE	Lys
463	ATC	CAC	ACG	GUG	GGT.	GGC	GIC	AIG	ACG	100	CIG	AIC	DAM
	asn	thr	thr	ile	pro	thr	lys	lys	ser	gln	ile	phe	ser
	AAC												
477									_			_	_
	tyr												
ACG 491	TAC	GCG	GAC	AAC	CAG	CCG	GGC	GIG	CAC	ATC	CAG	GIC	TTT
	gly	alu	aro	ala	met	thr	lvs	asp	CVS	his	leu	leu	alv
	GGG												
515	-												
	phe												
ACA	TTC	GAC	CTG	TCC	GGC	ATC	CCG	CCG	GCG	CCG	CGC	GGT	GIG

FIGURE 1-4





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FIGURE 1-5

641 asn gln glu ala ser lys glu glu tyr glu his arg gln lys AAC CAG GAG GCC AGC AAG GAG GAG TAC GAG CAC CGC CAG AAG 655 glu leu glu asn leu cys thr pro ile met thr lys met tyr GAG CTG GAG AAC CTG TGC ACG CCC ATC ATG ACG AAG ATG TAC 669 gln gly met gly ala gly gly met pro gly gly met pro CAG GGC ATG GGC GGC GGC GGT ATG CCC GGA GGT ATG CCT 683 gly gly met pro gly gly met pro gly gly ala asn pro ser GGT GGA ATG CCC GGG GGC ATG CCT GGT GGC GCG AAC CCG TCG 697 ser ser ser gly pro lys val glu glu val asp OP TCT TCG TCA GGA CCG AAG GTG GAG GAA GTG GAC TGA GAGCGCATCC CTGAAGATGTTCCCATGGCGCGTCTGCTCGCGAACGAATAACCCGTTGGTTTTCTCC CTTGTAGAGCGTAGAGGTCTGCGACAAACCCAGCCGCCATCACTATTTTTATTATTGG GTATTGTCATTGCGATGGCACTTGTGCTGTTGAGGGCACCACGGTTGCCTCTGCCATT TTTGTTGCTGACTGACGCCTGTGTGCGTCTCCTTGTACCGCCGGCTTCCTTTCCTCCT TTCTCCCCCCCCTCCTTCCCCCTGT

CLUSTERED PAIR-WISE ALIGNMENT listed in clustered order, in 'identity (no translation)' alphabet of:

1.	Mhyhsp70	(1-600)	7.	x17 0	(1-647)
2.	Bmehsp70	(1-605)	8.	humhsp70	(1-641)
3.	dnaK	(1-638)	9.	chkhsp70	(1-635)
4.	tc70kd	(1-669)	10.	mzehsp70	(1-646)
6.	rathsp70	(1-646)	11.	smahsp70	(1-620)

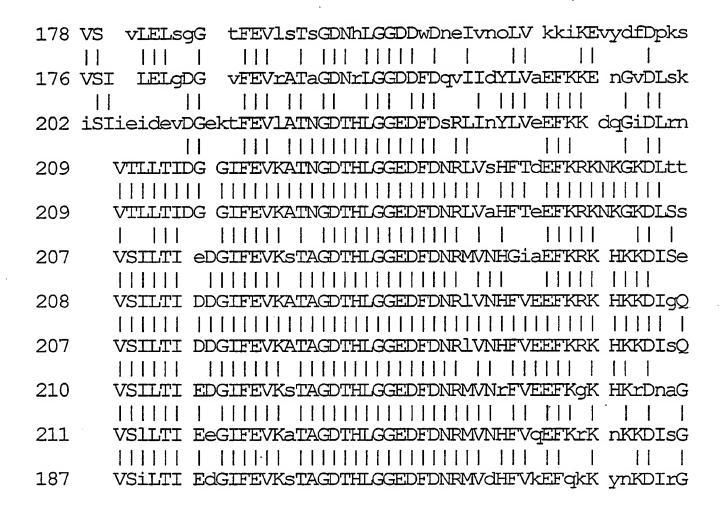
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1	MSKII GIDLGTINSCVA	vlEGgePkV	ipNPEGnRTTPS	VVAFKNGErqV
		1 1. 1	1 11 11111	
1	MgKII GIDLGTINSCVA	imdGttPRV	leNaEGdRTTPS:	iiaytqDGEtLV
1	MIYEGAI GIDLGTTYSCV	GVWQNERVE	IIANDQGNRTTPS	YVAFTDtERLI
1	MIYEGAI GIDLGTTYSCV	GVWQNERVE	IIANDQGNRTTPS	YVAFTDsERLI
1	MskGpA VGIDLGTTYSCV		IIANDQGNRTTPS	
1	MATKGVA VGIDLGTTYSCV	GVFQHGKVE:	IIANDQGNRTTPS	YVAFTDTERLI
1	MA KaaA VGIDLGTTYSCV	GVFQHGKVE:	IIANDQGNRTTPS	YVAFTDTERLI
1	msgkGPAIGIDLGTTYSCV		IIANDQGNRTTPS	
		1		
1	makseGPAIGIDLGTTYSCV		IIANDQGNRTTPS	•
_		• • •		
1		fOHakVE:	IIANDOGNRTTPS	YVartidserta

52	GdaAKRQ1eTNP ealaSiKR	JIMG.
		111
50	GevAKRQAiTNP NTIiSvKR	hMG
51	GqpAKRQAvTNPqNTlFaiKRLIGRrFqDeeVQrDvsimPFKiia	adnGD
		11
52	GDAAKNQVAMNPTNIVFDAKRLIGRKFSDpVVQSDMKHWPFK V	'iTKGDDKP
52	GDAAKNQVAMVPTNIVFDAKRLIGRKFSDsVVQSDMKHWPFK V	VIKGDDKP
		1
52	GDAAKNQVAMNPINIVFDAKRLIGRrFdDaVVQSDMKHWPF mV	V nDaGrP
		1 1 1
53	GDAAKNQVAMNPQNIVFDAKRLIGRKFnDPVVQcDlKHWPF QV	V sDeGKP
52	GDAAKNQVAlnPQNIVFDAKRLIGRKFGDPVVQcDlKHWPF QV	' iNDGdKP
53	GDAAKNQVAMNPTNTifDAKRLIGRKydDPtVQSDMKHWPF RV	vNeGgKP
		1 11
54	GDAAKNQVAMNPTNTVFDAKRLIGRRFssPaVQSsMKlWP sr	hlglGdKP
32	GDGAKNOVAMNPTNTVFDALRIJGRRFDDPSVOSDMKNWP fe	vt.avGaK1

75	IDktV	rAne	erdY	iPe	Eis	Ak.	ILa	yLŁ	eY.	AE	ki	Gh	kVI	.KAV	ΙΊ	VP.	AYF	'dn/	1gR
	11 1	1]					- 11	- 1	11		1							
74.	TDhkVE	Aec	gKqY	tPQ	EmS	Ai	ILG	hLi	ζgΥ	AE	ZYS	Œ	[V2	KAV	/ΙΊ	VP.	AYF	ND	AeR
	11			11	1	1	1						111				111		
101	awVE	vkg	ąKma	pPQ	is	¡Ae	ΛLk	KM	kt.	AEc	IYE	Œ	DV.	'eA\	/ΙΊ	VP.	AYF	ND	AQR
			1		l						11	-		1					
104	VIQVQE	RGET	CKTE	NPE	Evs	SM	⁄Ls	KMF	ŒĹ	AES	SYL	GK	QV k	KAY	ΝΊ	VP.	AYF	NDS	SQR
												11		11			111		
104	VIQVQF																		
			11 1	11	1 1	11			1			11	1						
103	KVQVEY	KGET	CKSF	УPE	Evs	SM	ΛI	KMF	ŒĹ	AEZ	YYL	GK	LVI	NAV.	ΝΊ	VP.	AYF	NDS	5QR
	11 111		111	11												11			
104	KVkVEY	KGE	KSF	'fPE	EIS	SM	ΙI	KMF	Et	AEZ	YYL	Ghl	5AJ	NAV.	ΠI	VP.	AYF	NDS	SQR
				-11	111				1							11			
103	KVQVsY	KGEt	KaF	ype	EIS	SM	ΠI	KMF	EL	AEZ	YYL	GyI	PVI	<i>IAM</i> :	/ΙΊ	VP.	AYF	NDS	SQR
				11								1				11			
104	KVQVeY																		
105	mIvfnY		-											7An	ΤVĪ	VP.	AYF	NDS	SQR
			1 1	1					1			1	1	11					
82	kIcveY	KGE	KMF	SpE	EIS	SM	/Lt	KMK	Evi	AES	YL	Gr.	ľvs	/Ab	7iT	VP/	AYF	NDS	SOR

128	eATKNAGKLAGLQVERLINEPTAAALATGL	aK	Tekemk\	LVYD.	LGGG	IFD
		1				
126	QATKDAGKIAGLEVERIINEPTAAALAYGL (еK	TdedqTV	LVYD:	LGGG	IFD
		1	-			Ш
152	QATKDAGrIAGLEV kRIINEPTAAALAYGLD	K	gtgnRTi	aVYD	LGGG.	IFD
			1	i		
157	QATKDAGTIAGMEVLRIINEPTAAAIAYGLD	Kve	dGKERNV	LIFD	LGGG!	IFD
157	QATKDAGTIAGLEVLRIINEPTAAAIAYGLD	Kac	JeGKERNV	LIFD	LGGG:	IFD
		1	1 1111	1111		
156	QATKDAGTIAGLNVLRIINEPTAAAIAYGLD	Κk	vGaERNV	LIFD	LGGG.	IFD
		1	1 11			
157	QATKDAGV1AGLN1LR1INEPTAAA1AYGLD	Κç	garGEqNV	LIFD	LGGG.	ΓFD
156	QATKDAGVIAGLNVLRIINEPTAAAIAYGLD	rI	.gkGErNV	LIFD	LGGG.	ŒD
		1				
157	QATKDAGTITGINVMRIINEPTAAAIAYGID	KKgI	PraGEKNV	LIFD	GGG.	ŒD
		$\Pi^{-}\Pi$				
158	QATKDAGVIAGINVMRIINEPTAAAIAYGIDE	KKaT	SSGEKNV	LIFDI	LGGG"	TFD
135	OATKDAGaTAGINVIRTINEPTAAATAYGIDE	KK	VACETINI	ומידד. ד	GGGT	(।स

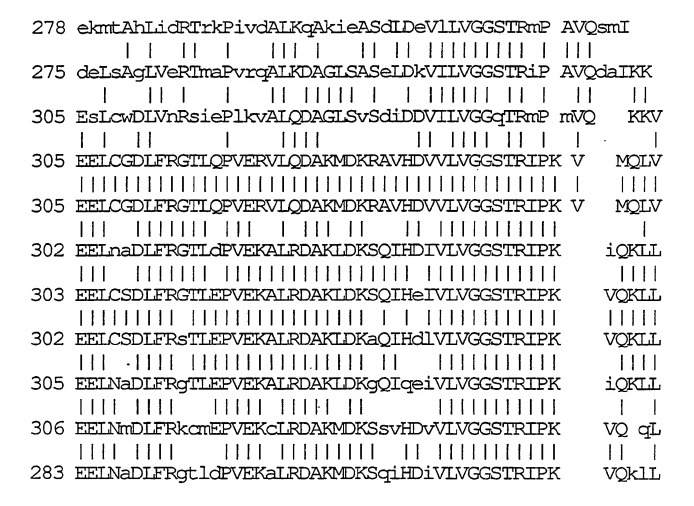
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227	DKMALTRLKeeAEKTKINLSI	JOSARI.	ASTELTAMP	CUCLI	verr.	LKKSer
225	DKMALQRLKdAAEKAKkdLS	gvtST	ŢĹSLPFITAG	eaGP	lHLEvs	SLSRAKF
255	DplamQRLKeAAEKAKieLS	SAqqTo	dvnLPyITAD?	ALGP	kHmnil	cvTRAKl
		11	1 1	1		
259	sqRALRRLRTACERAKRTLS	SA	AQATIEIDA	A LFOI	WDFQA:	TITRARF
259	NIRALRRIRTACERAKRTIS	SA	AQATIEIDA	A LFEN	VIDFQA	TITRARF
				1 1	11 .	
256	NKRAVRRLRTACERAKRTLS	S	StQASIEID	SLyE	SIDFYTS	SITRARE
		1				
257	NKRALRRLRTACdRAKRTLS	S	SsQASIEID	SLFEC	SIDFYT	ITRARF
256	NKRAVRRLRTACERAKRTLS	S	STGASLEID	SLFEC	SIDFYTS	SITRARE
		1 .				
257	NKRAVRRLRTACERATRILS	S	STQASIEID	SLFE	SIDFYTS	SITRARF
		1		1111		1111
258	NORAL RRL RTACERAKRILS	S	tAQTtIEID	SLFE	SIDFtpi	rssRARF
		1		• •		
236	NKRALRRLRTACERAKRTLS	S	sAQTnlEID	SLCC	SIDFyty	vitRARF

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326	ehtlnkkPnrsiNPDEVVAiGAAIQGGVLaG	eisDVlLLDVTPLtLGIE
•		
326	etggdPhKgVNPDEVVAlGAAIQGGVLIG	DVLDVVLLDVTPLSLGIE
353	aeffG KEPrKdVNPDEAVAiGAAVQGGVLTG	DVKDVLLLDVTPLSLGTE
354	SDFFGGKELNKSINPDEAVAYGAAVQAFILTGGKSF	
353	SDFFGGKELNKSINPDEA YGAAVQAFILTGGKSF	(Ölegiriri) vabrilete
351	QDFFNGKELNKSINPDEAVAYGAAVQAAILsGDKSF	CNVODITITIONATATE
352	QDFFNGRELNKSINPDEAVAYGAAVQAAIIMGDKSE	
250		
350	QDFFNGROLNKSINPDEAVGYGAAVQAAILMGDKSE	
254		
354	QDFFNGKELNKSINPDEAVAYGAAVQAAILMGDKSE	
25/	QDFFNGKELCKSINPDEAVAYGAAVQAAILSG egr	
334	UPFINGRELICASINEDEAVAIGAAVQAALIDG EGI	
221	QDFFNGKELnKSINPDEAVAYGAAVQAAILSGdkce	
22T	ONE LINGUEDING THE DEWAY TOWN OF WITHOUTH	

375	TIGGiaTpLIpRNTTIPvtKSQiFSTAeDnQTeVtIsVvQGERqiaADNKmL
392	TMGGVfTkLieRNTTIPTsKSQVFSTAaDsQTAVdIHVLQGERpmsADNKtL
422	TMGGVMItLIakNTTIPTKhSQVFSTAeDNQsAVtIHVLQGERkraADNKsL
426	TAGGVMTSLIKRNTTIPTKKSQIFSTYaDNQPGVHIQVFEGERaMIKDCHLL
425	TAGGVMTaLIKRNTTIPTKKSQIFSTYSDNQPGVHIQVFEGERtMFKDCHLL
424	TAGGVMIVLIKRNITIPTKQIQtFTTYSDNQPGVLIQVYEGERAMIKDNNLL
425	TAGGVMIVLIKRNITIPTKQIQsFTTYSDNQPGVLIQVfEGERAMIKDNNLL
424	TAGGVMTALIKRNSTIPTKQIQiFTTYSDNQPGVLIQVYEGERAMIKDNNLL
427	TAGGVMTALIKRNTTIPTKQIQLFTTYSDNQssVLvQVYEGERAMIKDNNLL
426	TAGGVMIVLIpRNITIPTKkeQvFsTYSDNQPGVLIQVYEGERArTKDNNLL
404	TAGGVMTaLIkRNTTIPTKqtQtftTYSDNQPGVLIQVfEGERAlTKDNNLL



427	GRENLSGIEAAPRGIPQIEVSESIDVNGITTVSAKDKRIGK EQTIT
425	GRFqLtdIpPAPRGvPQIEVSFDIDkNGIvnVrAKDlgTnK EQaIT
455	GqFnLdGInPAPRGmPQIEVTFDIDAdGILhVSAKDKnsGK EQkIT
459	GTFDLSGIPPAPRGVPQIEVTFDLDANGILnVSAEEKGTGKRNQIVIT
458	GTFDLSGIPPAPRGVPQIEVTFDLDANGILsVSAEEKGIGKRNQIVIT
457	GKFELtGIPPAPRGVPQIEVTFDIDANGILNVSAVdKSTGKeNKETIT
458	GKFELSGIPPAPRGVPQIEVTFDIDANGILNVSAVeKSsGKqNKITIT
456	GrFELSGIPPAP GVPQIEVTFDIDANGILNVtAtDKSTGKaNKITIT
460	GKFOLTGIPPAPRGVPQIEVTFDIDANGILNVSAvDKSTGKeNKITIT
459	GKFELSGIPPAPRGVPQItVTFDIDvNnILNVSAeDKtTGqkNKTTIT
437	GKFELSGIPPAPRGtPQIeVTFDIDaNgILNVSAvDKgTGkqNKITIT

473	IK	ntST LSeeEI	nkMiqEA	EENreAD	alKkdK
471	IK	SSTGLSdDEI	drMVkEA	EENAdad	KqRK
	11				11
501	IKA	SS GLneDEI	QkMVrDA	EaNAeAD	RK
	11	İ	1 11		
505	NDKGRLSKADIERMV	SeAAKYEsqDKe	QrerIDA	KNGL	ENYAFSv
		1 11111 11	111	1111	
507	NDKGRLSKADIERMV	Sdaakyeaedk	ahvIDA	KNGL	ENYAFSM
		1 11 111		11 1	
505	NDKFRLSKEDIERMV	QEAEKYKAED	ekQl	RdkVssKNsLl	ESYAFNM
			1		
506	NDKFRLSKEDIEKMV	QEAEKYKAdD	daQI	RERVďAKNALI	ESYAFNL
506	NDKGRLSKEeIERMV	QEAEKYKAED	EvQI	RERVSAKNALI	ESYAFNM
508	NDKGRLSKddIdRMV(<u>D</u> EAEKYKAED	EanI	RdRVgAKNsLI	ESYTYNM
	11111111				
507	NDKGRLSKEEIEkMV	<u> DEAEKYKAED</u>	Eev	kkVdAKNali	EnYaYNM
			1		
585	NDKGRLSKEEIErMV	adAdKYKAED	Ekgi	drVsAKNsLI	E syvyt



202	iettvraegiinQL	EKSITUQGEK	ıdpkqkellekq
501	EE VelRNeadQLv	fttEKtLkDlegKVEEA	evtkanea
		1111	
530	fEElVqtRN	qqdhlLhstrkqVEEA	gdklpaddKtaiEsaltaL
		1111	
552	KNIVNePNVAGK	ieeADKNtiTsAVEEALq	WLNnNQEASKEEYEHRQKEL
549	KNTINDPNVAGK	ldDADKNavTtAVEEALr	WLNdNQEASLEEYnHRQKEL
		1 11	
550	KaTVEDEklqGKI	nDEDKqkIldKCnEiIS	WLdkNQtaekeefeHQQKEL
551	KSmVEDEnvKGKI	SDEDKrtiseKCtqVIS	WLenNQLAEKEEyafQQKdL
552	KSaVEDEgLKGKI	SeaDKkKVLDKCQEVIS	WLDaNtlaekdefehkrkel
556	KqtVEDEkLKGKI	SdqDKqKVLDKCQEVIS	SLDrNQmAEKEEYEHKqKEL
	11		
555	rntikddKIask	:lpaeDKkKiEDavdgaISV	WLDsNQlaeveefeckmkel
		111	
529	mkaaveaelkeKIpes	sdhaviisKcED t.TSV	VLDvhOsAFkhFvesKreFT

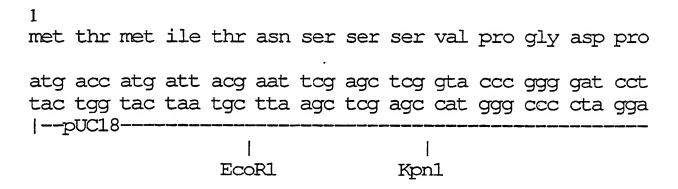


541	iqeLK	DL	lked	ktDEL	kLkldqie	eaaaqsfAQa
	11	11				
539	kdALKaaie	eknDLeeIkAK		kDELgei	vqaLtvKL	yeqAQ
	111		•		_	
573	EtALK	geDkaaIeAKN	1	qEL	aqvsqKLme	eiaqqqhAQ
						-
602	E	nlCtPImtKN	YQY	EMGaGGgm	PG	GMPgG
	1.		11		11	111 1
599	E	gVCaPIlsKM	YQX	EMG GGdg1	PG	GMPeG
					1	
600	E	KVCnPIITK	LYQ	Sa	a G	GMPGG
	1					1 111
601	E	KVCqPIITK	LYQ		G	GvPGG
						11
599	E	qVCNPII	sgLYQ		GAG	PG
	1	1111				
603	E	klCNPI	vtkLYQ		GAG	
	1	1111			111 .	
602	E	giCNPI		Iakmyxge	eGAG	
					1	
579	E	kvCaPI		I tkdvyc	nagG	



574 taqqa	ntsEsdpkaDDsntiDAEikqd
1111	
578 QAQQA G	EqgAqnDD VVDAEFEEVndDKK
609 Q QtA G A	daSAnnakdDD VVDAEFEEV kDKK
630 MPGGMPGG A	nPssssgpkwrkwteSASlkmfpwrrllanE
626 MPGGMPGG 1	mPG G mgggmGGaaASSGPkvEEVD
621 MPGG	fPG GGA ppsGG ASSGPTIEEVD
620 vPGG	mPGsscGAQarqGG nSGPTIEEVD
111	
618 PGG	fGAQgpkGG SGSGPTIEEVDO
620 aGA	GG SG GPTIEEVDO
1	1 11 11 11 11 11 11
620 MG	AaaGM dedapsGG SGaGPkIEEVDO
1	
596 M	pgGMheasgagGG SGkGPtIEEVD





29
ARG ASP LEU GLY PRO ASP ARG CYS ARG GLY ASP ILE ALA ARG
CGA GAT CTC GGG CCC GAT CGA TGC CGC GGC GAT ATC GCT CGA
GCT CTA GAG CCC GGG CTA GCT ACG GCG CCG CTA TAG CGA GCT

| | | Xho1

Position of pUC18 conserved sequences, addition endpoints and predicted partial amino acid sequence of the betagalactosidase fusion protein produced in pWHA148. A portion of the nucleotide sequence of pUC18 is designated

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FIGURE 4-2

by lower case letters; the nucleotide sequence of the pWHA148 synthetic oligonucleotide addition is designated by upper case letters. Numbers refer to the order of the espected amino acid sequence.



FIGURE 5-1

Translation of M. hyopneumoniae 74.5kD Antigen Gene

met ala lys glu ile ile leu gly ile asp leu gly thr thr ATG GCA AAA GAA ATC ATT TTA GGA ATC GAC CTT GGA ACA ACA 15 asn ser val val ala ile ile glu asn gln lys pro val val AAC TCA GTT GTT GCA ATT ATT GAA AAT CAA AAA CCT GTC GTT 30 leu glu asn pro asn gly lys arg thr thr pro ser val val CTC GAA AAT CCC AAC GGA AAA AGA ACA ACT CCA TCC GTT GTC ala phe lys asn asn glu glu ile val gly asp ala ala lys GCT TTT AAA AAC AAT GAA GAA ATT GTC GGG GAT GCA GCT AAA മ arg gln leu glu thr asn pro glu ala ile ala ser ile lys AGA CAA CTT GAA ACT AAC CCA GAA GCA ATC GCT TCA ATT AAA 75 arg leu met gly thr asp lys thr val arg ala asn glu arg AGA TIA ATG GGA ACT GAT AAA ACA GTI CGT GCA AAT GAA AGA 90 asp tyr ile pro glu glu ile ser ala lys ile leu ala tyr GAT TAT ATT CCT GAA GAA ATC TCG GCA AAA ATT CTT GCT TAT 105 leu lys glu tyr ala glu lys lys ile gly his lys val thr TTA AAA GAA TAT GCT GAG AAA AAG ATT GGT CAT AAA GTA ACA 120 lys ala val ile thr val pro ala tyr phe asp asn ala gln AAA GCA GTA ATT ACA GTA CCT GCT TAT TTT GAC AAT GCC CAA 135 arg glu ala thr lys asn ala gly lys ile ala gly leu gln CGT GAG GCA ACA AAA AAT GCC GGA AAA ATC GCT GGA TTA CAA 150 val glu arg ile ile asn glu pro thr ala ala ala leu ala GTA GAA AGA ATT ATA AAT GAA CCA ACA GCG GCC GCA CTT GCT 165 phe gly leu asp lys thr glu lys glu met lys val leu val TIT GGC CTT GAT AAA ACT GAA AAA GAA ATG AAA GTT CTT GTC



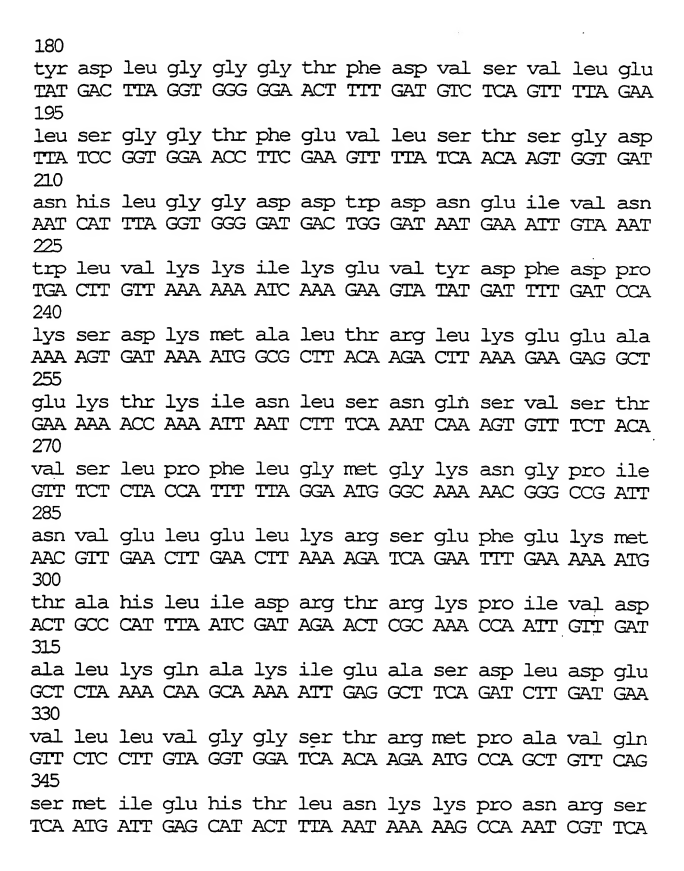


FIGURE 5-3

360 ile asn pro asp glu val val ala ile gly ala ala ile gln ATT AAT CCT GAT GAG GTA GTC GCA ATT GGT GCT GCA ATT CAA 375 gly gly val leu ala gly glu ile ser asp val leu leu leu GGG GGG GTT CTA GCT GGA GAG ATC AGT GAT GTT CTA CTT TTA 390 asp val thr pro leu thr leu gly ile glu thr leu gly gly GAT GIT ACT CCT ITA ACT ITA GGA ATT GAA ACT ITA GGI GGA 405 ile ala thr pro leu ile pro arg asn thr thr ile pro val ATT GCA ACA CCT TTG ATT CCA AGA AAT ACA ACA ATT CCG GTA 420 thr lys ser gln ile phe ser thr ala glu asp asn gln thr ACA AAA TCA CAA ATT TTC TCA ACA GCT GAG GAT AAT CAA ACC 435 glu val thr ile ser val val gln gly glu arg gln leu ala GAA GTA ACA ATT TCT GTT GTC CAA GGT GAA CGT CAA CTT GCA 450 ala asp asn lys met leu gly arg phe asn leu ser gly ile GCG GAT AAA AAG TIA GGT CGC ITT AAT ITA TCA GGA ATT 465 glu ala ala pro arg gly leu pro gln ile glu val ser phe GAA GCT GCT CCA CGA GGT CTT CCC CAG ATT GAA GTT AGT TTT 480 ser ile asp val asn gly ile thr thr val ser ala lys asp TCA ATT GAT GTC AAC GGG ATT ACA ACG GTT TCA GCA AAA GAT 495 lys lys thr gly lys glu gln thr ile thr ile lys asn thr AAA AAA ACC GGC AAA GAA CAA ACA ATT ACA ATT AAA AAT ACT 510 ser thr leu ser glu glu glu ile asn lys met ile gln glu TCA ACT TTA TCA GAA GAA GAA ATT AAT AAG ATG ATT CAG GAA 525 ala glu glu asn arg glu ala asp ala leu lys lys asp lys GCC GAA GAA AAT CGT GAA GCT GAT GCT CTT AAA AAA GAC AAA



FIGURE 5-4

540													
	_				_		_					gln	
ATC	GAG	ACA	ACA	GTT	CGT	GCC	GAA	GGG	CTT	ATT	TAA	CAA	CTT
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glu	lys	ser	ile	thr	asp	gln	gly	glu	lys	ile	asp	pro	lys
GAG	AAA	TCA	ATA	ACT	GAT	CAA	GGT	GAA	AAA	ATT	GAT	CCA	AAA
570													
gln	lys	glu	leu	leu	glu	lys	gln	ile	gln	glu	leu	lys	asp
CAA	AAA	GAA	TTA	CIT	GAA	AAA	CAA	ATT	CAA	GAA	TTA	AAA	GAT
585													
leu	leu	lys	glu	asp	lys	thr	asp	glu	leu	lys	leu	lys	leu
CTT	CTA	AAA	GAA	GAT	AAA	ACT	GAC	GAA	TTA	AAA	TTA	AAA	TTA
600													
asp	gln	ile	glu	ala	ala	ala	gln	ser	phe	ala	gln	ala	thr
GAC	CAA	ATT	GAA	GCA	GCT	GCC	CAA	TCT	TTT	GCG	CAG	GCA	ACC
ഖ5													
ala	gln	gln	ala	asn	thr	ser	glu	ser	asp	pro	lys	ala	asp
GCG	CAG	CAA	GCA	AAT	ACA	TCT	GAA	TCT	GAT	CCA	AAA	GCT	GAT
ഒ0													
asp	ser	asn	thr	ile	asp	ala	glu	ile	lys	gln	asp	∞	
GAT	TCA	AAC	ACA	TTA	GAT	GCT	GAA	ATC	AAG	CAG	GAT	TAA	

FIGURE 11-1

Translation of M. gallisepticum 67 kD Antigen Gene

1 met ser asn asn gly leu ile ile gly ile asp leu gly ATG TCT AAT AAT GGA TTA ATT ATT GGA ATT GAT CTT GGT 15 thr thr asn ser cys val ser val met glu gly ala gln lys ACC ACC AAC TCT TGT GTG TCT GTA ATG GAA GGT GCA CAA AAA 30 val val ile glu asn pro glu gly lys arg thr thr pro ser GTA GTA ATT GAA AAC CCA GAA GGT AAA AGA ACT ACT CCA TCA 45 val val ser tyr lys asn gly glu ile ile val gly asp ala GTA GTT TCA TAC AAA AAC GGT GAA ATT ATT GTT GGT GAT GCT മ ala lys arg gln met leu thr asn pro asn thr ile val ser GCT AAG CGT CAA ATG CTA ACT AAC CCA AAC ACT ATT GTT TCT *7*5 ile lys arg leu met gly thr ser lys lys val lys ile asn ATT AAG CGT TTA ATG GGA ACA AGT AAA AAA GTT AAG ATT AAT 90 asp lys gly val glu lys glu leu thr pro glu glu val ser GAC AAA GGT GIA GAA AAA GAA CTT ACT CCA GAA GAA GTT TCT 105 ala ser ile leu ser tyr leu lys asp tyr ala glu lys lys GCT AGC ATC TTA AGT TAT CTT AAA GAT TAC GCT GAA AAG AAA 120 thr gly gln lys ile ser arg ala val ile thr val pro ala ACT GGT CAA AAG ATT TCA AGA GCT GTA ATT ACT GTT CCA GCT 135 tyr phe asn asp ala glu arg gln ala thr lys thr ala gly TAC TTC AAC GAC GCT GAA CGT CAA GCT ACT AAA ACT GCT GGT 150 lys ile ala gly leu thr val glu arg ile ile asn glu pro AAG ATT GCT GGT TTA ACT GTA GAA AGA ATT ATT AAC GAA CCT



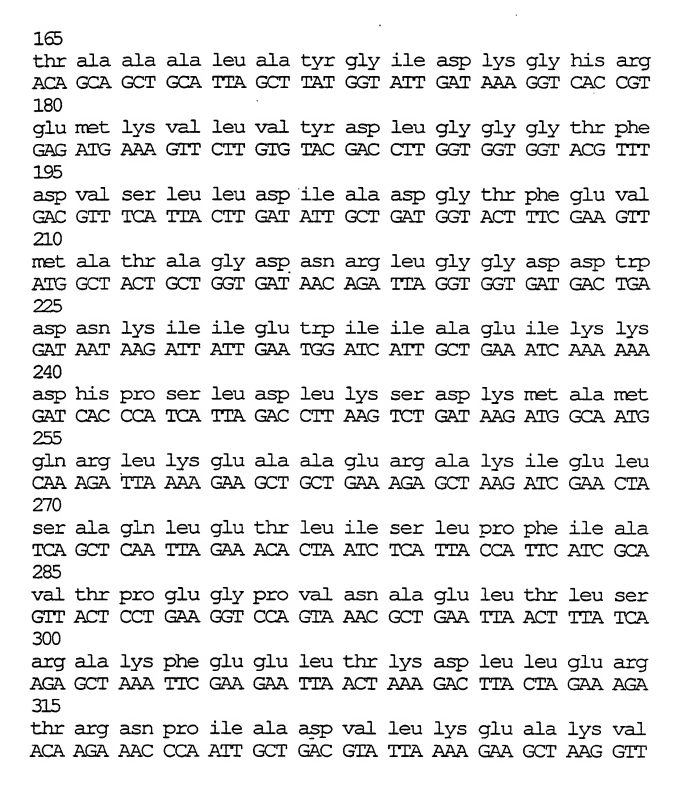


FIGURE 11-3

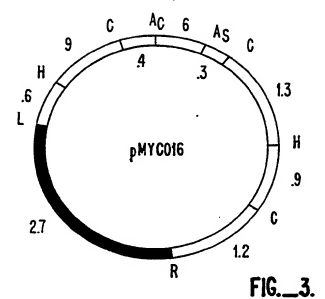
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GAT	CCT	AGT	CAA	GTT	GAT	GAA	ATT	CTT	TTA	GTA	GGT	GGT	TCT
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	AGA	AIG	CCT	GÇA	GTA	CAA	AAA	TTA	GTT	GAA	TCA	ATG	ATT
360		-	_									_	_
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	AAT	AAA	GCA	CCA	AAC	CGT	ACG	ATT.	AAC	CCT	GAC	GAA	GIA
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450													
	gln		_					_	_		_		
	CAA	GGT	GAA	CGT	CCA	ATG	GCT	AGA	GAA	AAC	AAA	TCA	TTA
465		,		_	,	7		-		-			-
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FIGURE 11-4

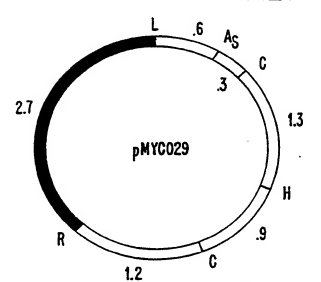
495 ile leu asn val lys ala lys asp leu thr thr gly lys glu ATT TTA AAT GIT AAG GCT AAA GAC TTA ACA ACT GGT AAA GAA 510 asn ser ile thr ile ser asn ser ser glu leu asp glu asn AAC AGT ATT ACG ATC TCT AAC TCA AGT GAA TTG GAT GAA AAC 525 glu ile gln arg met ile arg asp ala glu ala asn lys glu GAA ATC CAA AGA ATG ATC CGT GAT GCT GAA GCT AAC AAA GAA 540 arg asp ala ile val lys gln arg ile glu met arg tyr glu CGT GAC GCA ATC GTT AAA CAA AGA ATC GAA ATG CGT TAT GAA 555 gly glu gly ile val asn thr ile asn glu ile leu gly ser GGT GAA GGA ATT GTT AAT ACA ATT AAC GAA ATC CIT GGT TCT 570 lys glu ala glu ala leu pro ala gln glu lys ala ser leu AAA GAA GCA GAA GCG CTA CCT GCT CAA GAA AAA GCT AGC CTT 585 thr lys ile val asp gly ile asn gly ala leu lys ala glu ACT AAG ATC GIT GAT GGA ATT AAC GGT GCT CTT AAA GCT GAA 600 lys trp asp glu leu lys glu gln ile asp gly phe lys lys AAA TGA GAT GAA CTT AAA GAA CAG ATC GAC GGC TTC AAG AAA 615 trp arg asp met ser lys lys tyr gly gly glu ala TGA CGT GAT GAC ATG TCT AAG AAA TAC GGT GGT GGC GAA GCT 630 pro ala glu pro lys AM CCA GCC GAA CCT AAA TAG





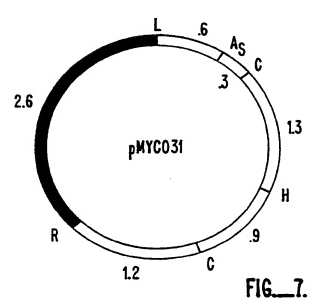
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- AC AccI
- As AsuII
- C ClaI
- H HindIII
- L HindIII-ClaI-PstI-AccI-EcoRI
- R ECORI
- pWHA148
- ☐ MYCOPLASMA



LEGEND

- As AsuII
- C ClaI
- H HindIII
- L HindII-ClaI-PstI
- R ECORI
- pWHA148
- **MYCOPLASMA**

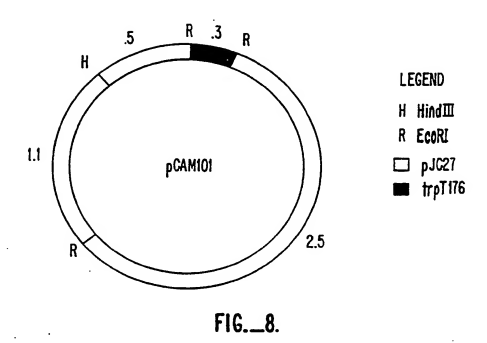


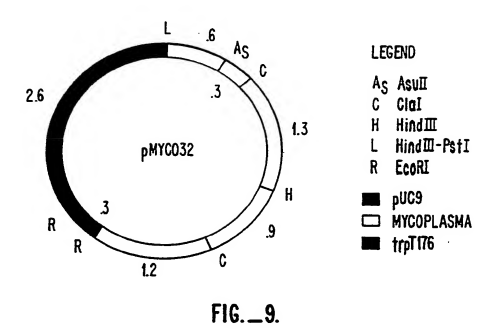
LEGEND

- As ASUII
- C CIQI
- H HindII
- L HindIII-PstI
- R EcoRI
- pUC9
- ☐ MYCOPLASMA

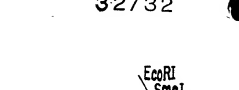
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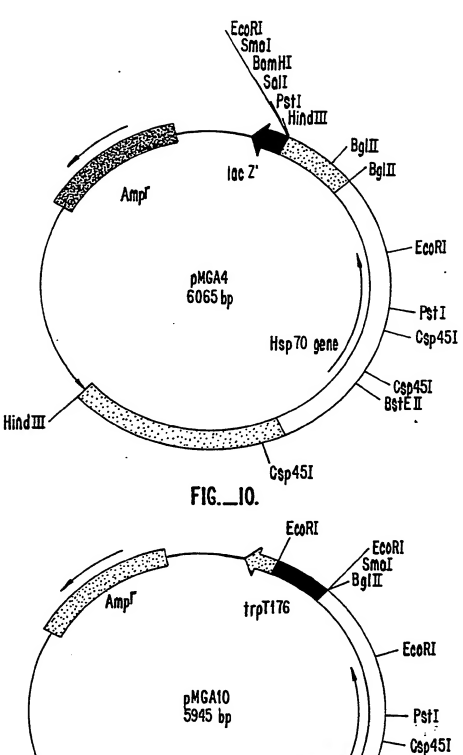
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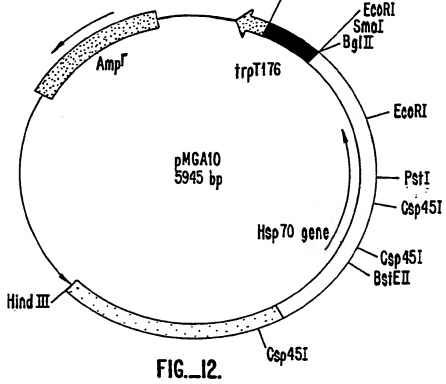




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International Application No. PCT/US89/03955

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